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(54) MELANGES HERBICIDES A EFFET SYNERGIQUE

(54) HERBICIDAL MIXTURES HAVING A SYNERGISTIC EFFECT

(57) L'invention concerne un mélange herbicide à effet synergique contenant A) au moins un dérivé benzoyle à substitution 3-hétérocyclyle de la formule (I), dans laquelle les variables ont la signification suivante: R¹, R³ sont hydrogène, halogène, alkyle, halogénure d'alkyle, alcoxy, halogénure d'alcoxy, alkylthio, alkylsulfinyle ou alkylsulfonyle, R² est un radical hétérocyclique choisi dans le groupe: thiazol-2-yl, thiazol-4-yl, thiazol-5-yl, isoxazol-3-yl, isoxazol-4-yl, isoxazol-5-yl, dihydroisoxazol-3-yl, 4,5-dihydroisoxazol-4-yl et 4.5dihydroisoxazol-5-yl, ceux-ci pouvant être substitués éventuellement une fois ou plusieurs fois par halogène, alkyle, alcoxy, halogénure d'alkyle, halogénure d'alcoxy ou alkylthio: R⁴ est hydrogène, halogène ou alkyle; R⁵ est alkyle; R⁶ est hydrogène ou alkyle; ou bien un de ses

(57) The invention relates to synergistic herbicidal mixtures containing A) at least one 3-heterocyclyl-substituted benzoyl derivative of formula (I) in which the variables have the following meaning: R¹, R³ represent hydrogen, halogen, alkyl, alkyl halide, alkoxy, alkoxy halide, alkylthio, alkyl sulfinyl, or alkyl sulfonyl; R² represents a heterocyclic radical selected from the group: thiazole-2-yl, thiazole-4-yl, thiazole-5-yl, isoxazol-3-yl, isoxazol-4-yl, isoxazol-5-yl, 4,5-dihydroisoxazol-3-yl, 4,5-dihydroisoxazol-5-yl, whereby these can be optionally substituted one time or a multiple number of times by halogen, alkyl, alkoxy, alkyl halide, alkoxy halide, alkylthio; R⁴ represents hydrogen, halogen or alkyl; R⁵ represents alkyl; R⁶ represents hydrogen or alkyl; or one of the

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sels écophiles, et B) une quantité produisant un effet synergique d'au moins un composé herbicide du groupe des inhibiteurs de l'acétyl CoA carboxylase (ACC), des inhibiteurs de l'acétolactate synthase (ALS), des amides, des herbicides de l'auxine, des inhibiteurs de transport de l'auxine, des inhibiteurs de biosynthèse de la carotinoïde, des inhibiteurs d'énolpyruvyl-shikimat-3-phosphatesynthase (ESPS), des inhibiteurs de la glutaminesynthétase, des inhibiteurs de la biosynthèse lipidique, des inhibiteurs de la mitose, des inhibiteurs de la protophorphyrinogèn-IX-oxydase, des inhibiteurs de la photosynthèse, des agents synergiques, des substances de croissance, des inhibiteurs de biosynthèse de paroi cellulaire, et de différents autres herbicides. L'invention concerne des agents contenant ces mélanges, ainsi que leur procédé de préparation et leur utilisation pour lutter contre des végétaux parasites.

environmentally compatible salts thereof; and B) a synergistically effective quantity of at least one herbicidal compound from the group of acetyl CoA carboxylase inhibitors (ACC), acetolactate synthase inhibitors (ALS), amides, auxin herbicides, auxin transport inhibitors, carotinoid biosynthesis inhibitors, enolpyruvyl-shikimat-3-phosphate synthase inhibitors (ESPS), glutamine synthetase inhibitors, biosynthesis inhibitors. mitosis inhibitors, protophorphyrinogen-IX-oxidase inhibitors, photosynthesis inhibitors, synergistic agents, growth substances, cell wall biosynthesis inhibitors and various other herbicides. The invention also relates to agents which contain these mixtures, to methods for producing these agents, and to the use thereof for controlling unwanted plants.

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Mit internationalem Recherchenbericht. Vor Ablauf der für Änderungen der Ansprüche zugelassenen Frist; Veröffentlichung wird wiederholt falls Anderungen eintreffen.

(54) Title: HERBICIDAL MIXTURES HAVING A SYNÉRGISTIC EFFECT

(54) Bezeichnung: HERBIZIDE MISCHUNGEN MIT SYNERGISTISCHER WIRKUNG

(57) Abstract

The invention relates to synergistic herbicidal mixtures containing A) at least one 3-heterocyclyl-substituted benzoyl derivative of formula (I) in which the variables have the following meaning: R¹, R³ represent hydrogen, halogen, alkyl, alkyl halide, alkoxy, alkoxy halide, alkylthio, alkyl sulfinyl, or alkyl sulfonyl; R2 represents a heterocyclic radical selected from the group: thiazole-2-yl, thiazole-4-yl, thiazole-5-yl, isoxazol-3-yl, isoxazol-4-yl, isoxazol-5-yl, 4,5-dihydroisoxazol-3-yl, 4,5-dihydroisoxazol-4-yl and 4,5-dihydroisoxazol-5-yl, whereby these can be optionally substituted one time or a multiple number of times by halogen, alkyl, alkoxy. alkyl halide, alkoxy halide, alkylthio; R4 represents hydrogen, halogen or alkyl; R5 represents alkyl; R6 represents hydrogen or alkyl; or one of the environmentally compatible salts thereof; and B) a synergistically effective quantity of at least one herbicidal compound from the group of acetyl CoA carboxylase inhibitors (ACC), acetolactate synthase inhibitors (ALS), amides, auxin herbicides, auxin transport inhibitors, carotinoid biosynthesis inhibitors, enolpyruvyl-shikimat-3-phosphate synthase inhibitors (ESPS), glutamine synthetase inhibitors, lipid biosynthesis inhibitors, mitosis inhibitors, protophorphyrinogen-IX-oxidase inhibitors, photosynthesis inhibitors, synergistic agents, growth substances, cell wall biosynthesis inhibitors and various other herbicides. The invention also relates to agents which contain these mixtures, to methods for producing these agents, and to the use thereof for controlling unwanted plants.

HERBICIDAL MIXTURES HAVING A SYNERGISTIC EFFECT

The present invention relates to a synergistic herbicidal mixture comprising

A) at least one 3-heterocyclyl-substituted benzoyl derivative of the formula I

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in which the variables have the following meanings:

 R^1 , R^3 are hydrogen, halogen, C_1 - C_6 -alkyl, C_1 - C_6 -haloalkyl, C_1 - C_6 -alkoxy, C_1 - C_6 -alkylsulfinyl or C_1 - C_6 -alkylsulfonyl;

is a heterocyclic radical selected from the group:
thiazol-2-yl, thiazol-4-yl, thiazol-5-yl,
isoxazol-3-yl, isoxazol-4-yl, isoxazol-5-yl,
4,5-dihydroisoxazol-3-yl, 4,5-dihydroisoxazol-4-yl
and 4,5-dihydroisoxazol-5-yl, it being possible for
the nine radicals mentioned to be unsubstituted or
mono- or polysubstituted by halogen, C₁-C₄-alkyl,
C₁-C₄-alkoxy, C₁-C₄-haloalkyl, C₁-C₄-haloalkoxy or
C₁-C₄-alkylthio;

R⁴ is hydrogen, halogen or C₁-C₆-alkyl;

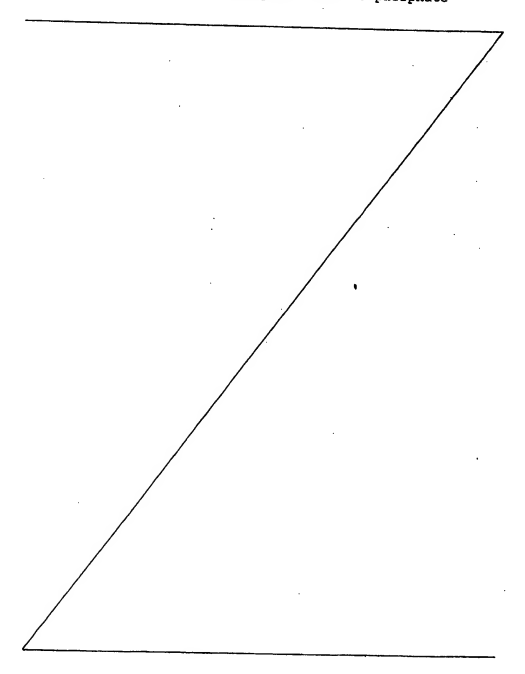
 R^5 is C_1-C_6 -alkyl;

R⁶ is hydrogen or C₁-C₆-alkyl;

or one of its environmentally compatible salts;

40 and

a synergistically effective amount of at least one herbicidal compound from the group of the acetyl-CoA carboxylase inhibitors (ACC), acetolactate synthase inhibitors (ALS), amides, auxin herbicides, auxin transport inhibitors, carotenoid biosynthesis inhibitors, enolpyruvylshikimate 3-phosphate



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synthase inhibitors (ESPS), glutamine synthetase inhibitors, lipid biosynthesis inhibitors, mitosis inhibitors, protoporphyrinogen IX oxidase inhibitors, photosynthesis inhibitors, synergists, growth substances, cell wall biosynthesis inhibitors and a variety of other herbicides.

The invention furthermore relates to herbicidal compositions comprising a herbicidally active amount of a synergistic herbicidal mixture as defined above and at least one liquid and/or solid carrier and, if desired, at least one surfactant.

Moreover, the invention relates to processes for the preparation of these compositions and to a method of controlling undesirable vegetation.

In crop protection products, it is always desirable to increase the specific activity of an active ingredient and the reliability 20 of action. It is an object of the present invention to increase the activity of known, herbicidally active 3-heterocyclyl-substituted benzoyl derivatives of the formula I.

It is an object of the present invention to increase the 25 selective herbicidal activity of the 3-heterocyclyl substituted benzoyl derivatives of the formula I against undesirable harmful plants.

We have found that this object is achieved by the mixtures defined at the outset. We have furthermore found herbicidal compositions which comprise these mixtures, processes for their preparation, and methods of controlling undesirable vegetation. In the last-mentioned cases, it is irrelevant whether the herbicidally active compounds of the components A) and B) are formulated and applied jointly or separately and in which sequence they are applied in the case of separate application.

The mixtures according to the invention show a synergistic effect; the compatibility of the herbicidally active compounds of components A) and B) for certain crop plants is generally retained.

Suitable components B are, as acetyl-CoA carboxylase inhibitors 45 (ACC), for example, cyclohexenone oxime ethers, phenoxyphenoxypropionic esters or arylaminopropionic acids. The acetolactate synthase inhibitors (ALS) include, inter alia,

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imidazolinones, pyrimidyl ethers, sulfonamides or sulfonyl ureas. Relevant auxin herbicides are, inter alia, pyridine carboxylic acids, 2,4-D or benazolin. Lipid biosynthesis inhibitors which are used are, inter alia, anilides, chloroacetanilides,

- 5 thioureas, benfuresate or perfluidone. Suitable mitosis inhibitors are, inter alia, carbamates, dinitroanilines, pyridines, butamifos, chlorthal-dimethyl (DCPA) or maleic hydrazide. Examples of protoporphyrinogen IX oxidase inhibitors are, inter alia, diphenyl ethers, oxadiazoles, cyclic imides or
- 10 pyrazoles. Suitable photosynthesis inhibitors are, inter alia, propanil, pyridate, pyridafol, benzothiadiazinones, dinitrophenols, dipyridylenes, ureas, phenols, chloridazon, triazine, triazinone, uracils or biscarbamates. The synergists are, inter alia, oxiranes. Examples of suitable growth substances
- 15 are aryloxyalkanoic acids, benzoic acids or quinolinecarboxylic acids. The group "various other herbicides" is to be understood as meaning, inter alia, the classes of the active ingredients dicloropropionic acids, dihydrobenzofurans, phenylacetic acids and individual herbicides mentioned below whose mechanism of 20 action is not (fully) understood.

Other suitable components B are active compounds selected from the group of the amides, auxin transport inhibitors, carotenoic biosynthesis inhibitors, enolpyruvylshikimate 3-phosphate

25 synthase inhibitors (EPSPS), glutamine synthetase inhibitors and cell wall synthesis inhibitors.

Examples of herbicides which can be used in combination with the 3-heterocyclyl-substituted benzoyl derivatives of formula I according to the present invention are, inter alia:

- B1 acetyl-CoA carboxylase inhibitors (ACC), for example
 - cyclohexenone oxime ethers, such as alloxydim, clethodim, cloproxydim, cycloxydim, sethoxydim, tralkoxydim, butroxydim, clefoxydim or tepraloxydim;
- phenoxyphenoxypropionic esters, such as clodinafop-propargyl (and, if appropriate, cloquintocet), cyhalofop-butyl, diclofop-methyl, fenoxaprop-ethyl, fenoxaprop-P-ethyl, fenthiapropethyl, fluazifop-butyl, fluazifop-P-butyl, haloxyfop-ethoxyethyl, haloxyfop-methyl, haloxyfop-P-methyl, isoxapyrifop, propaquizafop, quizalofop-ethyl, quizalofop-P-ethyl or quizalofop-tefuryl; or
 - arylaminopropionic acids, such as

flamprop-methyl	OI	flamprop-isopropyl;
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	•	ramprop-methyr or rramprop-isopropyl;
	(B2	acetolactate synthase inhibitors (ALS), for example
		- imidazolinones, such as imazapyr, imazaquin,
5		imazamethabenz-methyl (imazame), imazamoc, imazapic,
		imazethapyr or imazamethapyr;
		- Dyrimidyl ethors such
		- pyrimidyl ethers, such as pyrithiobac-acid,
		pyrithiobac-sodium, bispyribac-
10		sodium, KIH-6127 or pyribenzoxym;
		- sulfonamides, such as florasulam, flumetsulam or
		metosulam; or
		- sulfonylureas, such as amidosulfuron, azimsulfuron,
		bensulfuron-methyl, chlorimuron-ethyl, chlorsulfuron
		cinosulturon, cyclosulfamuron,
15		ethametsulfuron-methyl, ethoxysulfuron.
		flazasulfuron, halosulfuron-methyl, imazosulfuron,
		metsulfuron-methyl, nicosulfuron,
	•	primisulfuron-methyl, prosulfuron,
		pyrazosulfuron-ethyl, rimsulfuron,
20		sulfometuron-methyl, thifensulfuron-methyl,
		triasulfuron, tribenuron-methyl,
		triflusulfuron-methyl,
		N-[[[4-methoxy-6-(trifluoromethyl)-
		1,3,5-triazin-2-yl]amino]carbonyl]-2-(trifluoro-
25		methyl)-benzenesulfonamide, sulfosulfuron or
		idosulfuron;
		idosdifuton;
	В3	amides, for example
	20	
30		allidochlor (CDAA), benzoylprop-ethyl, bromobutide,
•		chlorthiamid, diphenamid, etobenzanid (benzchlomet),
		fluthiamide, fosamin or monalide;
	(24	
	(B4	auxin herbicides, for example
3.5		pyridinecarboxylic acids, such as
35		clopyralid or picloram; or
	•	clopyralid or picloram; or 2,4-D or benazolin;
	B5 a	nuxin transport inhibitors, for example
	-	naptalame or diflufenzopyr;
40		**
	B6 ca	rotenoid biosynthesis inhibitors, for example
	_	benzofenap, clomazone (dimethazone), diflufenican,
		fluorochloridone, fluridone, pyrazolynate,
		pyrazoxyfen igovaflutala daranta
15		pyrazoxyfen, isoxaflutole, isoxachlortole,
		mesotrione, sulcotrione (chlormesulone),
		ketospiradox, flurtamone, norflurazon or amitrol;

	В7	enolpyruvylshikimate-3-phosphate synthase inhibitors (ESPS), for example
		- glyphosate or sulfosate;
5	B8	glutamine synthetase inhibitors, for example
•		 bilanafos (bialaphos) or glufosinate-ammonium;
	В9	<pre>lipid biosynthesis inhibitors, for example - anilides, such as anilofos or mefenacet;</pre>
10		
		- chloroacetanilides, such as dimethenamid,
	`	S-dimethenamid, acetochlor, alachlor, butachlor,
		butenachlor, diethatyl-ethyl, dimethachlor,
		metazachlor, metolachlor, S-metolachlor,
15		<pre>pretilachlor, propachlor, prynachlor, terbuchlor, thenylchlor or xylachlor;</pre>
		 thioureas, such as butylate, cycloate, di-allate,
		dimepiperate, EPTC, esprocarb, molinate, pebulate,
		prosulfocarb, thiobencarb (benthiocarb), tri-allate
		or vernolate; or
20		- benfuresate or perfluidone;
	B10	mitosis inhibitors, for example
		- carbamates, such as asulam, carbetamid, chlorpropham,
25		orbencarb, pronamid (propyzamid), propham or tiocarbazil;
		- dinitroanilines, such as benefin, butralin,
		dinitramin, ethalfluralin, fluchloralin, oryzalin,
		pendimethalin, prodiamine or trifluralin;
		- pyridines, such as dithiopyr or thiazopyr; or
30		- butamifos, chlorthal-dimethyl (DCPA) or maleic
		hydrazide;
	B11	protoporphyrinogen IX oxidase inhibitors, for example
		 diphenyl ethers, such as acifluorfen,
35		acifluorfen-sodium, aclonifen, bifenox,
		chlornitrofen (CNP), ethoxyfen, fluorodifen,
		fluoroglycofen-ethyl, fomesafen, furyloxyfen,
		lactofen, nitrofen, nitrofluorfen or oxyfluorfen;
		 oxadiazoles, such as oxadiargyl or oxadiazon;
10		 cyclic imides, such as azafenidin, butafenacil,
		carfentrazone-ethyl, cinidon-ethyl,
		flumiclorac-pentyl, flumioxazin, flumipropyn,
		flupropacil, fluthiacet-methyl, sulfentrazone or
		thidiazimin; or
15		 pyrazoles, such as ET-751, JV 485 or nipyraclofen;

B12 photosynthesis inhibitors, for example

		•
		- propanil, pyridate or pyridafol;
		- benzothiadiazinones, such as bentazone;
		 dinitrophenols, for example bromofenoxim, dinoseb, dinoseb-acetate, dinoterb or DNOC;
5		 dipyridylenes, such as cyperquat-chloride,
		difenzoquat-methylsulfate, diquat or
		paraquat-dichloride;
		- ureas, such as chlorbromuron, chlorotoluron,
10		difenoxuron, dimefuron, diuron, ethidimuron, fenuron
		fluometuron, isoproturon, isouron, linuron,
		methabenzthiazuron, methazole, metobenzuron, metoxuron, monolinuron, neburon, siduron or
		tebuthiuron;
	٠	- phenols, such as bromoxynil or ioxynil;
15		- chloridazon;
		 triazines, such as ametryn, atrazine, cyanazine,
		desmetryn, dimethamethryn, hexazinone, prometon,
		prometryn, propazine, simazine, simetryn, terbumeton
20		terbutryn, terbutylazine or trietazine;
20		- triazinones, such as metamitron or metribuzin;
		 uracils, such as bromacil, lenacil or terbacil; or biscarbamates, such as desmedipham or phenmedipham;
		phenmedipham;
	B13	synergists, for example
25		- oxiranes, such as tridiphane;
	B14	growth substances, for example
		- aryloxyalkanoic acids, such as 2,4-DB, clomeprop,
		dichlorprop, dichlorprop-P (2,4-DP-P), fluoroxypyr,
30		MCPA, MCPB, mecoprop, mecoprop-P or triclopyr;
		- benzoic acids, such as chloramben or dicamba; or
		 quinolinecarboxylic acids, such as quinclorac or quinmerac;
		datimet ac;
35	B15	cell wall synthesis inhibitors, for example
		- isoxaben or dichlobenil;
	B16	various other herbicides, for example
40		- dichloropropionic acids, such as dalapon;
40		- dihydrobenzofurans, such as ethofumesate;
		- phenylacetic acids, such as chlorfenac (fenac); or
		 aziprotryn, barban, bensulide, benzthiazuron, benzofluor, buminafos, buthidazole, buturon,
		cafenstrole, chlorbufam, chlorfenprop-methyl,
45		chloroxuron, cinmethylin, cumyluron, cycluron,
		cyprazine, cyprazole, dibenzyluron, dipropetryn,
		dymron, eglinazin-ethyl, endothall, ethiozin,

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flucabazone, fluorbentranil, flupoxam, isocarbamid, isopropalin, karbutilate, mefluidide, monuron, napropamide, napropanilide, nitralin, oxaciclomefone, phenisopham, piperophos, procyazine, profluralin, pyributicarb, secbumeton, sulfallate (CDEC), terbucarb, triaziflam, triazofenamid or trimeturon;

or their environmentally compatible salts.

- 10 Of particular importance are the following herbicides which can be used in combination with the 3-heterocyclyl-substituted benzoly [sic] derivatives of the formula I according to the present invention:
- 15 Bl acetyl-CoA carboxylase inhibitors (ACC), for example
 - cyclohexenone oxime ethers, such as alloxydim, clethodim, cloproxydim, cycloxydim, sethoxydim, tralkoxydim, butroxydim, clefoxydim or tepraloxydim;
- phenoxyphenoxypropionic esters, such as

 clodinafop-propargyl (and, if appropriate, cloquintocet),
 cyhalofop-butyl, diclofop-methyl, fenoxaprop-ethyl,
 fenoxaprop-P-ethyl, fenthiaprop-ethyl, fluazifop-butyl,
 fluazifop-P-butyl, haloxyfop-ethoxyethyl,
 haloxyfop-methyl, haloxyfop-P-methyl, isoxapyrifop,
 propaquizafop, quizalofop-ethyl, quizalofop-P-ethyl or
 quizalofop-tefuryl; or
 - arylaminopropionic acids, such as flamprop-methyl or flamprop-isopropyl;
- 30 B2 acetolactate synthase inhibitors (ALS), for example
 - imidazolinones, such as imazapyr, imazaquin, imazamethabenz-methyl (imazame), imazapic, imazethapyr or imazamethapyr;
- pyrimidyl ethers, such as pyrithiobac-acid,
 pyrithiobac-sodium, bispyribac-sodium, KIH-6127 or pyribenzoxym;
 - sulfonamides, such as flumetsulam or metosulam; or
 - sulfonylureas, such as amidosulfuron, azimsulfuron, bensulfuron-methyl, chlorimuron-ethyl, chlorsulfuron, cinosulfuron, cyclosulfamuron, ethametsulfuron-methyl, ethoxysulfuron, flazasulfuron, halosulfuron-methyl, imazosulfuron, metsulfuron-methyl, nicosulfuron, primisulfuron-methyl, prosulfuron, pyrazosulfuron-ethyl,
- rimsulfuron, sulfometuron-methyl, thifensulfuron-methyl, triasulfuron, tribenuron-methyl, triflusulfuron-methyl, N-[[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin-2-

yl]amino]carbonyl]-2-(trifluoromethyl)benzenesulfonamide, sulfosulfuron or idosulfuron;

- B3 amides, for example
- 5 allidochlor (CDAA), benzoylprop-ethyl, bromobutide, chlorthiamid, diphenamid, etobenzanid (benzchlomet), fluthiamide, fosamine or monalide;
 - B4 auxin herbicides, for example
- pyridinecarboxylic acids, such as clopyralid or picloram; or
 - 2,4-D or benazolin;
 - B5 auxin transport inhibitors, for example
- 15 naptalame or diflufenzopyr;
 - B6 carotenoid biosynthesis inhibitors, for example
 - benzofenap, clomazone (dimethazone), diflufenican, fluorochloridone, fluridone, pyrazolynate, pyrazoxyfen, isoxaflutole, isoxachlortole, mesotrione, sulcotrione (chlormesulone), flurtamone, norflurazon or amitrol;
 - B7 enolpyruvylshikimate 3-phosphate synthase inhibitors (ESPS), for example
- 25 glyphosate or sulfosate;
 - B8 glutamine synthetase inhibitors, for example
 - bilanafos (bialaphos) or glufosinate-ammonium;
- 30 anilides, such as anilofos or mefenacet;
 - chloracetanilides, such as dimethenamid, S-dimethenamid, acetochlor, alachlor, butachlor, butenachlor, diethatyl-ethyl, dimethachlor, metazachlor, metolachlor, S-metolachlor, pretilachlor, propachlor, prynachlor,
- 35 terbuchlor, thenylchlor or xylachlor;
 - thioureas, such as butylate, cycloate, di-allate, dimepiperate, EPTC, esprocarb, molinate, pebulate, prosulfocarb, thiobencarb (benthiocarb), tri-allate or vernolate; or
- 40 benfuresate or perfluidone;
 - B10 mitosis inhibitors, for example
- carbamates, such as asulam, carbetamide, chlorpropham, orbencarb, pronamide (propyzamide), propham or thiocarbazil;

- dinitroanilines, such as benefin, butralin, dinitramine, ethalfluralin, fluchloralin, oryzalin, pendimethalin, prodiamine or trifluralin;
- pyridines, such as dithiopyr or thiazopyr; or
- 5 butamifos, chlorthal-dimethyl (DCPA) or maleic hydrazide;
 - B11 protoporphyrinogen IX oxidase inhibitors, for example
 - diphenyl ethers, such as acifluorfen, acifluorfen-sodium, aclonifen, bifenox, chlornitrofen (CNP), ethoxyfen,
- fluorodifen, fluoroglycofen-ethyl, fomesafen, furyloxyfen, lactofen, nitrofen, nitrofluorfen or oxyfluorfen;
 - oxadiazoles, such as oxadiargyl or oxadiazon;
- cyclic imides, such as azafenidin, carfentrazone-ethyl,
 cinidon-ethyl, flumiclorac-pentyl, flumioxazin,
- flumipropyn, flupropacil, fluthiacet-methyl, sulfentrazone or thidiazimin; or
 - pyrazoles, such as ET-751, JV 485 or nipyraclofen;
- 20 B12 photosynthesis inhibitors, for example
 - propanil, pyridate;
 - benzothiadiazinones, such as bentazon;
 - dinitrophenols, such as bromofenoxim, dinoseb, dinoseb-acetate, dinoterb or DNOC;
- 25 dipyridylenes, such as cyperquat-chloride, difenzoquat-methylsulfate, diquat or paraquat-dichloride;
 - ureas, such as chlorbromuron, chlorotoluron, difenoxuron, dimefuron, diuron, ethidimuron, fenuron, fluometuron, isoproturon, isouron, linuron, methabenzthiazuron,
- methazole, metobenzuron, metoxuron, monolinuron, neburon, siduron or tebuthiuron;
 - phenols, such as bromoxynil or ioxynil;
 - chloridazon;
- triazines, such as ametryn, atrazine, cyanazine,
 desmetryn, dimethamethryn, hexazinone, prometon,
 prometryn, propazin, simazine, simetryn, terbumeton,
 terbutryn, terbutylazine or trietazine;
 - triazinones, such as metamitron or metribuzin;
 - uracils, such as bromacil, lenacil or terbacil; or
- biscarbamates, such as desmedipham or phenmedipham;
 - B13 synergists, for example
 - oxiranes, such as tridiphane;
- 45 B14 growth substances, for example

- aryloxyalkanoic acids, such as 2,4-DB, clomeprop, dichlorprop, dichlorprop-P (2,4-DP-P), fluoroxypyr, MCPA, MCPB, mecoprop, mecoprop-P or triclopyr;
- benzoic acids, such as chloramben or dicamba; or
- 5 quinolinecarboxylic acids, such as quinclorac or quinmerac;

10

B16 various other herbicides, for example

- dichloropropionic acids, such as dalapon;
- dihydrobenzofurans, such as ethofumesate;
- phenylacetic acids, such as chlorfenac (fenac); or
- aziprotryn, barban, bensulide, benzthiazuron, benzofluor, buminafos, buthidazole, buturon, cafenstrole, chlorbufam, chlorfenprop-methyl, chloroxuron, cinmethylin, cumyluron, cycluron, cyprazine, cyprazole, dibenzyluron, dipropetryn, dymron, eglinazin-ethyl, endothall, ethiozin, flucabazone, fluorbentranil, flupoxam, isocarbamid, isopropalin, karbutilate, mefluidide, monuron, napropamide, napropanilide, nitralin, oxaciclomefone, phenisopham, piperophos, procyazine, profluralin, pyributicarb, secbumeton, sulfallate (CDEC),

25 terbucarb, triazofenamid or trimeturon;

or their environmentally compatible salts.

The 3-heterocyclyl-substituted benzoyl derivatives of the formula 30 I are disclosed in WO 96/26206, WO 97/41116, WO 97/41117 and WO 97/41118.

They can exist, or be used, in the form of the pure enantiomers and also as racemates or diastereomer mixtures. The

- 35 3-heterocyclyl-substituted benzoyl derivatives of the formula I and the herbicidally active compounds from amongst groups B1 to B16 may also exist in the form of their environmentally compatible salts. Suitable salts are, in general, the salts of those cations, or the acid addition salts of those acids, whose
- 40 cations, or anions, respectively, do not adversely affect the herbicidal action of the active ingredients.

Suitable cations are, in particular, ions of the alkali metals, preferably lithium, sodium and potassium, of the alkaline earth metals, preferably calcium and magnesium, and of the transition metals, preferably manganese, copper, zinc and iron, and also ammonium, it being possible in this case, if desired, for one to

four hydrogen atoms to be replaced by C₁-C₄-alkyl, hydroxy-C₁-C₄-alkyl, C₁-C₄-alkoxy-C₁-C₄-alkyl, hydroxy-C₁-C₄-alkoxy-C₁-C₄-alkyl, phenyl or benzyl, preferably ammonium, dimethylammonium, diisopropylammonium, tetrabutylammonium, 2-(2-hydroxyeth-1-oxy)eth-1-yl ammonium, di(2-hydroxyeth-1-yl)ammonium, trimethylbenzylammonium, furthermore phosphonium ions, sulfonium ions, preferably tri(C₁-C₄-alkyl)sulfonium and sulfoxonium ions, preferably, 10 tri(C₁-C₄-alkyl)sulfoxonium.

Anions of suitable acid addition salts are mainly chloride, bromide, fluoride, hydrogen sulfate, sulfate, dihydrogen phosphate, hydrogen phosphate, nitrate, hydrogen carbonate, carbonate, hexafluorosilicate, hexafluorophosphate, benzoate and the anions of C₁-C₄-alkanoic acids, preferably formate, acetate, propionate and butyrate.

- The herbicidally active compounds from amongst groups B1 to B16 are described, for example, in
 - "Herbizide [Herbicides]", Hock, Fedtke, Schmidt, 1st edition, Thieme 1995 (s. "quinclorac" p. 238, "molinat" p. 32,
- "butachlor" p. 32, "pretilachlor" p. 32, "dithiopyr" p. 32, "mefenacet" p. 32, "fenoxapropethyl" p. 216, "dimepiperate" p. 32, "pyrazolynate" p. 146, "pyrazoxyfen" p. 146, "bensulfuronmethyl" p. 31, "pyrazosulfuron-ethyl" p. 31, "cinosulfuron" p. 31, "benfuresate" p. 233, "bromobutide"
- p. 243, "dymron" p. 243, "dimethyametryn" p. 118, "esprocarb" p. 229, "pyributicarb" p. 32, "cinemthylin" p. 32, "propanil" p. 32, "2,4-D" p. 30, "bentazon" p. 30, "azimsulfuron (DPX-A-8947)" p. 175, "mecoprop-P" p. 237, "chlorpropham" p. 205, "ethoxyfen" p. 30, "haloxyfop-P-methyl" p. 38,
- "haloxyfop-ethoxyethyl" p. 38, "flumiclorac-pentyl" p. 35, "flupropacil" p. 143, "nipyraclofen" p. 145, "metosulam" p. 33, "ethametsulfuron-methyl" p. 36, "thifensulfuron-methyl" p. 35, "pyrithiobac acid" p. 181);
- "Agricultural Chemicals", Book II Herbicides, 1993 (s.
 "thiobencarb" p. 85, "benzofenap" p. 221, "napropanilid"
 p. 49, "piperophos" p. 102, "anilofos" p. 241, "imazosulfuron
 (TH-913)" p. 150, "etobenzamid (HW-52)" p. 54, "sulcotrione
 (ICIA-0051)" p. 268, "poast" p. 253, "focus" p. 222,
- "dimethenamid" p. 48, "sulfosate" p. 236, "2,4-DB" p. 10,
 "dichlorprop-P" p. 6, "flupoxam" p. 44, "prosulfocarb" p. 84,
 "quinmerac" p. 233, "metazachlor" p. 64, "flurtamone" p. 265,

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"imazamethabenz-methyl" p. 153, "clodinafop-propargyl"
p. 214, "fenoxaprop-P-ethyl" p. 208, "fluazifop-P-butyl"
p. 207, "quizalofop-P-ethyl" p. 210, "quizalofop-terfuryl"
p. 211, "flumioxazin" p. 43, "flumipropyn" p. 267,
"sulfentrazone" p. 261, "thiazopyr" p. 226,
"pyrithiobac-sodium" p. 266, "flumetsulam" p. 227,
"amidosulfuron" p. 151, "halosulfuron-methyl" p. 148,
"rimsulfuron" p. 138, "tribenuron-methyl" p. 139,
"triflusulfuron-methyl" p. 137, "primisulfuron-methyl"
p. 147);
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- "Agricultural Chemicals", Book II Herbicides, 13th Edition (s.
 "carfenstole" p. 284, "sulfosulfuron" p. 145,
 "ethoxysulfuron" p. 149, "pyribenzoxym" p. 279,
 "diflufenzopyr" p. 90, "ET-751" p. 278, "carfentrazone-ethyl"
 p. 267, "fluthiacet-methyl" p. 277, "imazapic" p. 160,
 "butenachlor" p. 54, "tiocarbazil" p. 84, "fluthiamide"
 p. 62, "isoxaflutole" p. 283, "butroxydim" p. 259,)
- "Short Review of Herbicides & PGRs 1991, Hodogaya Chemicals (s. "furyloxyfen" p. 142, "triazofenamid" p. 268, "thenylchlorid (NSK-850)" p. 52, "cumyluron (JC-940)" p. 90, "pendimethalin (AC-92553)" p. 58, "buthidazole" p. 88, 25 "cyprazole" p. 38, "allidochlor" p. 48, "benzoylprop-ethyl" p. 38, "chlorthiamid" p. 150, "diphenamid" p. 34, "flamprop-methyl" p. 40, "fosamin" p. 232, "isoxaben" p. 42, "monalide" p. 32, "naptalam" p. 36, "pronamid" p. 34, "bialaphos" p. 234, "glufosinate-ammonium" p. 234, 30 "glyphosate" p. 232, "amitrol" p. 254, "clomeprop p. 20, "dichlorprop" p. 6, "fenoprop" p. 8, "fluroxypyr" p. 156, "MCPA" p. 4, "MCPB" p. 8, "mecoprop" p. 6, "napropamide" p. 16, "triclopyr" p. 154, "chloramben" p. 28, "dicamba" p. 26, "clomazone" p. 268, "diflufenican" p. 42, 35 "fluorochloridone" p. 266, "fluridone" p. 156, "asulam" p. 112, "barban" p. 100, "butylate" p. 106, "carbetamide" p. 36, "chlorobufam" p. 100, "cycloate" p. 108, "desmedipham" p. 104, "di-allate" p. 106, "EPTC" p. 108, "orbencarb"
- p. 104, "di-allate" p. 106, "EPTC" p. 108, "orbencarb"
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 "xylachlor" p. 52, "alloxydim" p. 260, "clethodim" p. 270,
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       "metribuzin" p. 202, "bromacil" p. 180, "lenacil" p. 180,
       "terbacil" p. 180, "benazolin" p. 262, "bensulide" p. 228,
       "benzofluor" p. 266, "butamifos" p. 228, "DCPA" p. 28,
       "dichlobenil" p. 148, "endothal" p. 264, "mefluidide" p. 306,
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       "perfluidone" p. 260, "terbuchlor" p. 48);
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PCT/EP 96/03996.

- "Global Herbicide Directory" First Edition, 1994 (s. "oxadiargyl" p. 96);
- "European Directory of Agrochemical Products" Volume 2 Herbicides" Fourth Edition, (s. "buminafos" p. 255).

Moreover, the compound "DEH-112" is disclosed in European Patent Application EP-A 302 203. The compound "tepraloxydim" is described in DE-A 33 36 140; the compound "cinidon-ethyl" in DE-A 36 03 789 and the compound "fluorbentranil" in EP-A 84 893. Other compounds are known from "Brighton Crop Protection Conference - Weeds - 1993 (S. "thidiazimin" p. 29, "AC-322140" p. 41, "KIH-6127" p. 47, "prosulfuron" p. 53, "KIH-2023" p. 61, "metobenzuron" p. 67). The compound "carfenstrole (CH-900)" is mentioned in EP-A 332 133, and the compound N-[[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin-2-yl]amino]-carbonyl]-2-(trifluoromethylbenzenesulfonamide) is described in

The assignment of the active ingredients to the respective mechanisms of action is based on current knowledge. If several mechanisms of action apply to one active ingredient, this substance was only assigned to one mode of action.

Preferred with regard to the synergistic herbicidal action of the mixtures according to the invention are those 3-heterocyclyl-substituted benzoyl derivatives of the formula I in which the variables have the following meanings, either alone 30 or in combination:

- halogen, C₁-C₆-alkyl, C₁-C₆-alkylthio, C₁-C₆-alkylsulfinyl or C₁-C₆-alkylsulfonyl; especially preferably halogen, such as chlorine or bromine, C₁-C₆-alkyl, such as methyl or other as C₁-C₁-alkyl, such as methyl or other as C₁-C₂-alkyl, as methyl or other as C₁-C₂-alkyl, such as methyl or other as C₂-C₂-alkyl, as
- C₁-C₆-alkyl, such as methyl or ethyl, or C₁-C₆-alkylsulfonyl, such as methylsulfonyl or ethylsulfonyl; very particularly preferably chlorine, methyl or methylsulfonyl;
- a heterocyclic radical selected from the group:
 isoxazol-3-yl, isoxazol-5-yl and 4,5-dihydroisoxazol-3-yl, it
 being possible for the three radicals mentioned to be
 unsubstituted or mono- or polysubstituted by halogen,
 C1-C4-alkyl, C1-C4-alkoxy, C1-C4-haloalkyl, C1-C4-haloalkoxy or
 C1-C4-alkylthio;
 especially preferably isoxazol-5-yl, 3-methyl-isoxazol-5-yl

especially preferably isoxazol-5-yl, 3-methyl-isoxazol-5-yl, 4,5-dihydroisoxazol-3-yl, 5-methyl-4,5-dihydroisoxazol-yl, 5-ethyl-4,5-dihydroisoxazol-3-yl or

4,5-dimethyl-4,5-dihydroisoxazol-3-yl; also preferred is a heterocyclic radical selected from the group: thiazol-2-yl, thiazol-4-yl, thiazol-5-yl, isoxazol-4-yl, 4,5-dihydroisoxazol-4-yl and

- 4,5-dihydroisoxazol-5-yl, it being possible for the six radicals mentioned to be unsubstituted or mono- or polysubstituted by halogen, C₁-C₄-alkyl, C₁-C₄-alkoxy, C₁-C₄-haloalkyl, C₁-C₄-haloalkoxy or C₁-C₄-alkylthio;
- halogen, C₁-C₆-alkyl, C₁-C₆-alkylthio, C₁-C₆-alkylsulfinyl or C₁-C₆-alkylsulfonyl; especially preferably halogen, such as chlorine or bromine, C₁-C₆-alkylthio, such as methylthio or ethylthio, C₁-C₆-alkylsulfinyl, such as methylsulfinyl or ethylsulfinyl, or C₁-C₆-alkylsulfonyl, such as methylsulfonyl or ethylsulfonyl; very particularly preferably chlorine, methylsulfonyl or ethylsulfonyl;
- 20 R4 hydrogen or methyl; especially preferably hydrogen;
- is C₁-C₆-alkyl, such as methyl, ethyl, propyl, 1-methylethyl,
 butyl, 1-methylpropyl or 2-methylpropyl;
 especially preferably methyl, ethyl or 1-methylethyl;
- R⁶ hydrogen or C₁-C₆ alkyl, such as methyl or ethyl; especially preferably hydrogen or methyl.
 30

Very particularly preferred are those 3-heterocyclyl-substituted benzoyl derivatives of the formula Ia, in particular the compounds Ia.1 to Ia.53, which are mentioned in Table 1 which follows:

40

Table 1

5

R⁵, OH R¹

la

	No.	R ¹	R ²	R ³	R ⁴	R ⁵	R ⁶
10	la.1	CI	4,5-dihydroisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	CH ₃
	Ia.2	CI	4,5-dihydroisoxazol-3-yl	CI	Н	CH ₃	CH ₃
	la.3	CI	4,5-dihydroisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	Н
	Ia.4	CI	4,5-dihydro-5-methylisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	Н
	la.5	Cl	4,5-dihydro-5,5-dimethylisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	Н
15	la.6	Cl	4,5-dihydro-5-ethylisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	Н
	la.7	CI	4,5-dihydro-5,5-diethylisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	Н
	la.8	Cl	4,5-dihydro-5-chloromethylisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	Н
	Ia.9	CI	4,5-dihydroisoxazol-3-yl	SCH ₃	Н	CH ₃	Н
20	la.10	Cl	4,5-dihydro-5-ethoxyisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	Н
	Ia.11	Cl	4,5-dihydro-5-methoxyisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	Н
	Ia.12	CI	4,5-dihydro-4,5-dimethylisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	Н
	la.13	CI	4,5-dihydro-5-thioethylisoxazol-3-yl	SO ₂ CH ₃	Н	СН3	Н
	Ia.14	Cl	4,5-dihydro-5-trifluoromethylisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	Н
25	Ia.15	SCH ₃	4,5-dihydroisoxazol-3-yl	SCH ₃	Н	СН3	Н
	la.16	CI	4,5-dihydroisoxazol-3-yl	SO ₂ CH ₃	Н	C ₂ H ₅	Н
	Ia.17	CI	4,5-dihydroisoxazol-3-yl	CI	H	C ₂ H ₅	Н
	la.18	Cl.	4,5-dihydro-5-methylisoxazol-3-yl	SO ₂ CH ₃	Н	C ₂ H ₅	Н
30	la.19	CI	4,5-dihydro-5,5-dimethylisoxazol-3-yl	SO ₂ CH ₃	Н	C ₂ H ₅	Н
	la.20	Cl	4,5-dihydro-5-ethylisoxazol-3-yl	SO ₂ CH ₃	H	C ₂ H ₅	Н
	la.21	CI	4,5-dihydro-5,5-diethylisoxazol-3-yl	SO ₂ CH ₃	Н	C ₂ H ₅	Н
	Ia.22	Cl	4,5-dihydroisoxazol-3-yl	SCH ₃	Н	C ₂ H ₅	Н
	Ia.23	Cl	4,5-dihydro-5-chloromethylisoxazol-3-yl	SO ₂ CH ₃	Н	C ₂ H ₅	Н
35	la.24	CI	4,5-dihydroisoxazol-3-yl	SOCH ₃	Н	C ₂ H ₅	Н
	Ia.25	Cl	4,5-dihydro-5-ethoxyisoxazol-3-yl	SO ₂ CH ₃	Н	C ₂ H ₅	Н
	Ia.26	CI	4,5-dihydro-4,5-dimethylisoxazol-3-yl	SO ₂ CH ₃	Н	C ₂ H ₅	Н
	Ia.27	CI	4,5-dihydro-5-thioethylisoxazol-3-yl	SO ₂ CH ₃	Н	C ₂ H ₅	Н
40	Ia.28	Cl	4,5-dihydro-5-trifluoromethylisoxazol-3-yl	SO ₂ CH ₃	Н	C ₂ H ₅	Н
	la.29	SCH ₃	4,5-dihydroisoxazol-3-yl	SCH ₃	Н	C ₂ H ₅	Н
į	Ia.30	Cl	4,5-dihydroisoxazol-3-yl	SO ₂ CH ₃	Н	i-C ₄ H ₉	Н
	la.31	CH ₃	4,5-dihydroisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	CH ₃
	la.32	CH ₃	4,5-dihydroisoxazol-3-yl	CI	H	CH ₃	CH ₃
45	Ia.33	CH ₃	4,5-dihydroisoxazol-3-yl	SO ₂ CH ₃	H	CH ₃	Н
	la.34	CH ₃	4,5-dihydro-5-methylisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	Н
	Ia.35	CH ₃	4,5-dihydro-5,5-dimethylisoxazol-3-yl	SO ₂ CH ₃	H	CH ₃	Н

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45

	No.	R1	R ²	R ³	R ⁴	R ⁵	R ⁶
	la.36	CH ₃	4,5-dihydro-5-ethylisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	H
	la.37	CH ₃	4,5-dihydro-5,5-diethylisoxazol-3-yl	SO ₂ CH ₃	H	CH ₃	Н
5	Ia.38	CH ₃	4,5-dihydroisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	Н
_	la.39	CH ₃	4,5-dihydro-4,5-dimethylisoxazol-3-yl	SO ₂ CH ₃	Н	CH ₃	Н
	la.40	CH ₃	4,5-dihydroisoxazol-3-yl	SO ₂ CH ₃	H	C ₂ H ₅	Н
	la.41	CH ₃	4,5-dihydroisoxazol-3-yl	CI	H	C ₂ H ₅	Н
	Ia.42	CH ₃	4,5-dihydro-5-methylisoxazol-3-yl	SO ₂ CH ₃	H	C ₂ H ₅	Н
10	Ia.43	CH ₃	4,5-dihydro-5,5-dimethylisoxazol-3-yl	SO ₂ CH ₃	Н	C ₂ H ₅	Н
	Ia.44	CH ₃	4,5-dihydro-5-ethylisoxazol-3-yl	SO ₂ CH ₃	Н	C ₂ H ₅	Н
	Ia.45	CH ₃	4,5-dihydro-5-diethylisoxazol-3-yl	SO ₂ CH ₃	H	C ₂ H ₅	Н
	Ia.46	CH ₃	4,5-dihydro-4,5-dimethylisoxazol-3-yl	SO ₂ CH ₃	H	C ₂ H ₅	Н
15	Ia.47	CH ₃	4,5-dihydroisoxazol-3-yl	SO ₂ CH ₃	Н	i-C ₄ H ₉	Н -
	Ia.48	Cl	2-thiazolyl	SO ₂ CH ₃	Н	CH ₃	CH ₃
	Ia.49	CI	2-thiazolyl	SO ₂ CH ₃	Н	CH ₃	Ĥ
	Ia.50	Cl	2-thiazolyl	SO ₂ CH ₃	Н	C ₂ H ₅	Н
	la.51	CH ₃	2-thiazolyl	SO ₂ CH ₃	H	CH ₃	CH ₃
20	la.52	Cl	3-methylisoxazol-5-yl	SO ₂ CH ₃	Н	CH ₃	Н
	la.53	Cl	3-methylisoxazol-5-yl	SO ₂ CH ₃	H	C ₂ H ₅	Н

Also very particularly preferred are the compounds Ib, in particular the compounds 1b.1 to 1b.53, which differ from the compounds Ia.1 to Ia.53 only by the fact that they are present as the sodium salt:

• Also very particularly preferred are the compounds Ic, in particular the compounds Ic.1 to Ic.53, which differ from the compounds Ia.1 to Ia.53 only by the fact that they are present as the lithium salt:

 Also very particularly preferred are the compounds Id, in particular the compounds Id.1 to Id.53, which differ from the compounds Ia.1 to Ia.53 only by the fact that they are present as the potassium salt:

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$$\begin{array}{c|c}
R^6 & O & R^1 \\
N & O & R^2 \\
R^5 & O & R^4
\end{array}$$
Id

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Also very particularly preferred are the compounds Ie, in particular the compounds Ie.1 to Ie.53, which differ from the compounds Ia.1 to Ia.53 only by the fact that they are present as the ammonium salt:

Very particularly preferred are, especially, the compounds
 Ia, especially the compounds Ia.1 to Ia.53.

 Very particularly preferred are, moreover, the 3-heterocyclyl substituted benzoyl derivatives of the formula I where

is a heterocyclic radical selected from amongst the group:
thiazol-2-yl, thiazol-4-yl and thiazol-5-yl, it being
possible for the three radicals mentioned to be unsubstituted
or mono- or polysubstituted by halogen, C₁-C₄-alkyl,
C₁-C₄-alkoxy, C₁-C₄-haloalkyl, C₁-C₄-haloalkoxy or
C₁-C₄-alkylthio.

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Very particularly preferred are, especially, the 3-heterocyclyl-substituted benzoyl derivatives of the formula I, where ${\tt R^4}$ is hydrogen.

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Very particularly preferred are, moreover, the 3-heterocyclyl substituted benzoyl derivatives of the formula I where

R² is a heterocyclic radical selected from the group:

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isoxazol-3-yl, isoxazol-4-yl and isoxazol-5-yl, it being possible for the three radicals mentioned to be unsubstituted or mono- or polysubstituted by halogen, C_1-C_4 -alkyl, C_1-C_4 -alkoxy, C_1-C_4 -haloalkyl, C_1-C_4 -haloalkoxy or C_1-C_4 -alkylthio.

Very particularly preferred are, especially, the 3-heterocyclyl-substituted benzoyl derivatives of the formula I, where

- R² is isoxazol-3-yl which can be unsubstituted or mono- or polysubstituted by halogen, C₁-C₄-alkyl, C₁-C₄-alkoxy, C₁-C₄-haloalkyl, C₁-C₄-haloalkoxy or C₁-C₄-alkylthio.

 R⁴ is hydrogen.
- Very especially preferred are also in particular the 3-heterocyclyl-substituted benzoyl derivatives of the formula I where
- R² is isoxazol-5-yl, which can be unsubstituted or mono- or polysubstituted by halogen, C₁-C₄-alkyl, C₁-C₄-alkoxy, C₁-C₄-haloalkyl, C₁-C₄-haloalkoxy or C₁-C₄-alkylthio; R⁴ is hydrogen.
- Most particularly preferred is

 4-[2-chloro-3-(3-methyl-isoxazol-5-yl)-4-methylsulfonyl-benzoyl]-1-methyl-5-hydroxy-1H-pyrazole.
- Very particularly preferred are, moreover, the
 3-heterocyclyl-substituted benzoyl derivatives of the formula
 I where
- R² is a heterocyclic radical selected from the group:
 4,5-dihydroisoxazol-3-yl, 4,5-dihydroisoxazol-4-yl and
 4,5-dihydroisoxazol-5-yl, it being possible for the three
 radicals mentioned to be unsubstituted or mono- or
 polysubstituted by halogen, C₁-C₄-alkyl, C₁-C₄-alkoxy,
 C₁-C₄-haloalkyl, C₁-C₄-haloalkoxy or C₁-C₄-alkylthio.
- Very particularly preferred are, especially, the
 3-heterocyclyl-substituted benzoyl derivatives of the
 formula I where

- R² is 4,5-dihydroisoxazol-3-yl which can be unsubstituted or mono- or polysubstituted by halogen, C₁-C₄-alkyl, C₁-C₄-alkoxy, C₁-C₄-haloalkyl, C₁-C₄-haloalkoxy or C₁-C₄-alkylthio; and
- 5 R4 is hydrogen.

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Most particularly preferred are the 3-heterocyclyl-substituted benzoyl derivatives of the formula I where

- R1 is halogen or C1-C6-alkyl; and
- R^3 is C_1-C_6 -alkylsulfonyl.
- Most especially preferred is

 4-[2-chloro-3-(4,5-dihydroisoxazol-3-yl)-4-methylsulfonyl-benzoyl]-1-methyl-5-hydroxy-1H-pyrazole.

Most particularly preferred is also

4-[2-methyl-3-(4,5-dihydroisoxazol-3-yl)-4-methylsulfonyl-benzoyl]-1-methyl-5-hydroxy-1H-pyrazole.

With a view to the synergistic herbicidal action of the mixtures according to the invention, compounds from amongst groups B1 to B14, are preferred as component B).

In particular, compounds from amongst the classes of active ingredients mentioned below are preferred, or the following 30 compounds are very particularly preferred:

- Bl acetyl-CoA carboxylase inhibitors (ACC):
- cyclohexenone oxime ethers, in particular cycloxydim, sethoxydim or tralkoxydim, preferably sethoxydim or tralkoxydim; or
 - phenoxyphenoxypropionic esters, in particular clodinafop-propargyl (and, if appropriate, cloquintocet), fenoxaprop-ethyl or fenoxaprop-P-ethyl, preferably clodinafop-propargyl (and, if appropriate, cloquintocet) or fenoxaprop-p-ethyl [sic];
 - B2 acetolactate synthase inhibitors (ALS):
- imidazolinones, in particular imazapyr, imazaquin,
 imazamethabenz, imazethapyr or imazamoc, preferably
 imazapyr;
 - pyrimidyl ethers, in particular pyrithiobac sodium;

- sulfonamides, in particular florasulam, flumetsulam or metosulam, preferably metosulam; or
- sulfonylureas, in particular halosulfuron-methyl, nicosulfuron, primisulfuron-methyl, prosulfuron, rimsulfuron, thifensulfuron-methyl, tribenuron-methyl, N-[[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin-2-yl]amino]carbonyl]-2-(trifluoromethyl)-benzenesulfonamide or sulfosulfuron;

10 B3 amides:

- fluthiamide;
- B4 auxin herbicides:
 - pyridinecarboxylic acids, in particular clopyralid; or
- -15 2,4-D;
 - B5 auxin transport inhibitors:
 - diflufenzopyr;
 - B6 carotenoid biosynthesis inhibitors:
- 20 isoxaflutole, mesotrione, isoxachloride, ketospiradox or sulcotrione (chlormesulone), in particular isoxaflutole or sulcotrione;
- B7 enolpyruvylshikimate-3-phosphate synthase inhibitors (ESPS):
 25 glyphosate or sulfosate;
 - B8 glutamin synthetase inhibitors:
 - glufosinate-ammonium;
- 30 B9 lipid biosynthesis inhibitors:
 - chloroacetanilides, in particular dimethenamid,
 S-dimethenamid, acetochlor, metolachlor or S-metolachlor,
 - thioureas, in particular benthiocarb;
- 35 B10 mitosis inhibitors:
 - dinitroanilines, in particular pendimethalin;
 - Bll protoporphyrinogen IX oxidase inhibitors:
- diphenyl ethers, in particular acifluorfen or
 acifluorfen-sodium;
 - oxadiazoles, in particular oxadiargyl; or
 - cyclic imides, in particular butafenacil, carfentrazone-ethyl, cinidon-ethyl or flumiclorac-pentyl, preferably carfentrazone-ethyl, cinidon-ethyl or
- 45 flumidorac-pentyl;
 - pyrazoles, in particular JV 485;

B12 photosynthesis inhibitors:

- pyridate or pyridafol, in particular pyridate;
- benzothiadiazinones, in particular bentazone;
- dipyridylenes, in particular paraquat-dichloride;
- ureas, in particular diuron or isoproturon, preferably diuron;
 - phenols, in particular bromoxynil;
 - chloridazone;
 - triazines, in particular atrazine or terbutylazine; or
- 10 triazinones, in particular metribuzin;

Bl3 synergists:

oxiranes, in particular tridiphane;

15 Bl4 growth substances:

- aryloxyalkanoic acids, in particular fluoroxypyr, MCPA or mecoprop-P;
- benzoic acids, in particular dicamba; or
- quinolinecarboxylic acids, in particular quinclorac;

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B16 various other herbicides:

triaziflam.

Also preferred as component B) are compounds from amongst the groups B1, B2, B4 to B12 and B14.

In particular, compounds from amongst the classes of active ingredients mentioned below are preferred, or the following compounds are very particularly preferred:

- B1 acetyl-CoA carboxylase inhibitors (ACC):
 - cyclohexenone oxime ethers, in particular cycloxydim or sethoxydim;
- phenoxyphenoxypropionic esters, in particular clodinafop-propargyl (and, if appropriate, cloquintocet), fenoxaprop-ethyl or fenoxaprop-P-ethyl, preferably clodinafop-propargyl (and, if appropriate, cloquintocet);

40 B2 acetolactate synthase inhibitors (ALS):

- imidazolinones, in particular imazapyr, imazaquin, imazamethabenz or imazethapyr, preferably imazapyr;
- pyrimidyl ethers, in particular pyrithiobac-sodium;
- sulfonamides, in particular flumetsulam or metosulam,
 preferably metosulam; or

- sulfonylureas, in particular halosulfuron-methyl, nicosulfuron or N-[[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin-2-yl]amino]carbonyl]-2-(trifluoromethyl)benzenesulfonamide, preferably nicosulfuron or N-[[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin-2-yl]amino]carbonyl]-2-(trifluoromethyl)-benzenesulfonamide;
- B4 auxin herbicides:

- 2,4-D;

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- B5 auxin transport inhibitors:
 - diflufenzopyr;

B6 carotenoid biosynthesis inhibitors:

- 15 isoxaflutole or sulcotrione, preferably isoxaflutole;
- 20 B8 glutamine synthetase inhibitors:
 - glufosinate-ammonium;
 - B9 lipid biosynthesis inhibitors:
 - chloracetanilide, in particular dimethenamid, S-dimethenamid, acetochlor, metolachlor or S-metolachlor;
 - thioureas, in particular benthiocarb;

Bl0 mitosis inhibitors:

dinitroaniline, in particular pendimethalin;

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- Bll protoporphyrinogen IX oxidase inhibitors:
 - diphenyl ethers, in particular acifluorfen;
 - cyclic imides, in particular carfentrazone-ethyl or cinidon-ethyl, preferably carfentrazone-ethyl;

- Bl2 photosynthesis inhibitors:
 - pyridate;
 - benzothiadiazinones, in particular bentazone;
 - dipyridylenes, in particular paraquat-dichloride;
- 40 ureas, in particular diuron or isobroturon, preferably diuron;
 - phenols, in particular bromoxynil;
 - chloridazon;
 - triazines, in particular atrazine or terbutylazine; or
- 45 triazinones, in particular metribuzin;

B14 growth substances:

- aryloxyalkanoic acids, in particular MCPA;
- benzoic acids, in particular dicamba;
- quinolinecarboxylic acids, in particular quinclorac.

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The following embodiments are especially preferred with a view to the synergistic herbicidal action of the mixtures according to the invention:

- In a particular embodiment, the synergistic herbicidal mixture according to the invention comprises, as component A), a 3-heterocyclyl-substituted benzoyl derivative of the formula I, where
- is a heterocyclic radical selected from the group:
 isoxazol-3-yl, isoxazol-5-yl and
 4,5-dihydroisoxazol-3-yl, the three radicals mentioned
 being unsubstituted or mono- or polysubstituted by
 halogen, C₁-C₄-alkyl, C₁-C₄-alkoxy, C₁-C₄-haloalkyl,
 C₁-C₄-haloalkoxy or C₁-C₄-alkylthio;
- in particular isoxazol-5-yl, 3-methyl-isoxazol-5-yl, 4,5-dihydroisoxazol-3-yl, 5-methyl-4,5-dihydroisoxazol-3-yl, 5-ethyl-4,5-dihydroisoxazol-3-yl or 4,5-dimethyl-4,5-dihydroisoxazol-3-yl;
- and,
 as component B), at least one herbicidal compound from
 amongst the groups B1, B2, B4 to B12 and B14;
 in particular clodinafop (and, if appropriate, cloquintocet),
 diflufenzopyr, imazethapyr, flumetsulam, pyrithiobac-sodium,
 nicosulfuron, N-[[4-methoxy-6-(trifluoromethyl)-1,3,5triazin-2-yl]amino]-carbonyl]-2-(trifluoromethyl)benzenesulfonamide, clopyralid, 2,4-D, isoxaflutole, glyphosate,
 glufosinate-ammonium, dimethenamide, S-dimethenamide,
 acetochlor, metolachlor, S-metolachlor, pendimethalin,
- carfentrazone-ethyl, pyridate, bentazone, diuron, bromoxynil, atrazine, terbutylazine, metribuzine or dicamba.

Very particularly preferred are mixtures which comprise, as component A), 4-[2-chloro-3-(4,5-dihydroisoxazol-3-yl)-4-methylsulfonylbenzoyl]-1-methyl-5-hydroxy-1H-pyrazole.

Very particularly preferred are also mixtures which comprise, as component A), 4-[2-methyl-3-(4,5-dihydroisoxazol-3-yl)-4-methylsulfonylbenzoyl]-1-methyl-5-hydroxy-1H-pyrazole.

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Very particularly preferred are also mixtures which comprise, as component A), 4-[2-chloro-3-(3-methyl-isoxazol-5-yl)-4-methylsulfonylbenzoyl]-1-methyl-5-hydroxy-1H-pyrazole.

- 5 In another particular embodiment, the synergistic herbicidal mixture according to the invention comprises, as component A), a 3-heterocyclyl-substituted benzoyl derivative of the formula I where
- is a heterocyclic radical selected from the group:
 thiazol-2-yl, thiazol-4-yl, thiazol-5-yl, isoxazol-4-yl,
 4,5-dihydroisoxazol-4-yl and 4,5-dihydroisoxazol-5-yl, it
 being possible for the six radicals mentioned to be
 unsubstituted or mono- or polysubstituted by halogen,
 C1-C4-alkyl, C1-C4-alkoxy, C1-C4-haloalkyl,
- 15 C₁-C₄-haloalkoxy or C₁-C₄-alkylthio;
 and, as component B), at least one herbicidal compound from amongst the groups B1, B2, B4 to B12 and B14;
 in particular clodinafop (and, if appropriate, cloquintocet), diflufenzopyr, imazethapyr, flumetsulam, pyrithiobac-sodium, nicosulfuron, N-[[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin-2-yl]amino]-carbonyl]-2-(trifluoromethyl)benzene-sulfonamide, clopyralid, 2,4-D, isoxaflutole, glyphosate, glufosinate-ammonium, dimethenamide, S-dimethenamide, acetochlor, metolachlor, S-metolachlor, pendimethalin, carfentrazone-ethyl, pyridate, bentazone, diuron, bromoxynil,
 - In a further particular embodiment, the synergistic herbicidal mixture according to the invention comprises, as component A), a 3-heterocyclyl-substituted benzoyl derivative of the formula I where

atrazine, terbutylazine, metribuzine or dicamba.

R² is a heterocyclic radical selected from the group consisting of 4,5-dihydroisoxazol-3-yl, 4,5-dihydro-isoxazol-4-yl and 4,5-dihydroisoxazol-5-yl, where the three abovementioned radicals may be unsubstituted or mono- or polysubstituted by halogen, C₁-C₄-alkyl, C₁-C₄-alkoxy, C₁-C₄-haloalkyl, C₁-C₄-haloalkoxy or C₁-C₄-alkylthio;

and as component B) at least one herbicidal compound from amongst the groups B1, B2, B4 to B12 and B14;

- The synergistic herbicidal mixture according to the invention preferably comprises, as component B), at least one herbicidal compound from the following groups:

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	Bl	acetyl-CoA carboxylase inhibitors (ACC): cyclohexenone oxime ethers or phenoxypropionic esters;
5	В2	acetolactate synthase inhibitors (ALS): imidazolinones, pyrimidyl ethers, sulfonamides or sulfonylureas;
10	В4	auxin herbicides: pyridinecarboxylic acids or 2,4-D;
	. B5	auxin transport inhibitors;
15	В6	carotenoid biosynthesis inhibitors;
	В7	enolpyruvylshikimate 3-phosphate synthase inhibitors;
20	В8	glutamine synthetase inhibitors;
	В9	lipid biosynthesis inhibitors: chloroacetanilides or thioureas,
25	B10	mitosis inhibitors: dinitroanilines;
	B11	protoporphyrinogen IX oxidase inhibitors: diphenyl ethers, oxadiazoles, cyclic imides or pyrazoles;
30	B12	photosynthesis inhibitors: pyridate, pyridafol, benzothiadiazinone, dipyridylene, ureas, phenols, chloridazon, triazines
35		or triazinones, in particular pyridate, benzothinediazinone, dipyridylenes, ureas, phenols, chloridazon, triazines or triazinones;
40	B14	growth substances: aryloxyalkanoic acids, benzoic acids or quinolinecarboxylic acids.
	acco	earticular, the synergistic herbicidal mixture ording to the invention comprises, as component B), a tone herbicidal compound from the group:
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cycloxydim, sethoxydim, clodinafop (and, if appropriate, cloquintocet), fenoxaprop-ethyl, fenoxaprop-P-ethyl, imazapyr, imazaquin, imazamethabenz, imazethapyr, pyrithiobac-sodium, metosulam, halosulfuron-methyl, 5 nicosulfuron, N-[[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin-2-yl]amino]carbonyl]-2-(trifluoromethyl)benzenesulfonamide, flufenacet, 2,4-D, diflufenzopyr, isoxaflutole, sulcotrione, glyphosate, glufosinate-ammonium, dimethenamid, S-metolachlor, 10 benthiocarb, pendimethalin, acifluorfen, carfentrazone-ethyl, cinidon-ethyl, pyridate, bentazon, paraquat-dichloride, diuron, isoproturon, bromoxynil, chloridazon, atrazine, metribuzin, MCPA, dicamba and quinclorac. 15 Also preferably, the synergistic herbicidal mixture according to the invention comprises, as component B), at least one herbicidal compound from amongst the groups B1, B2, B4 to B11 and B14; 20 In particular, the synergistic herbicidal mixture according to the invention comprises at least one herbicidal compound from amongst the following groups: 25 Bl acetyl-CoA carboxylase inhibitors (ACC): cyclohexenone oxime ethers or phenoxypropionic esters; acetolactate synthase inhibitors (ALS): 30 imidazolinones, pyrimidyl ethers, sulfonamides or sulfonylureas; B4 auxin herbicides: 2,4-D; 35 **B**5 auxin transport inhibitors; **B6** carotenoid biosynthesis inhibitors; 40 **B**7 enolpyruvylshikimate 3-phosphate synthase inhibitors; glutamine synthetase inhibitors; B8 45 В9 lipid biosynthesis inhibitors:

chloroacetanilides or thioureas,

B10	mitosis	inhibitors
	dinitro	nilines;

- Bll protoporphyrinogen IX oxidase inhibitors: diphenyl ethers, oxadiazoles, cyclic imides or pyrazoles;
 - B14 growth substances: aryloxyalkanoic acid, benzoic acids or quinolinecarboxylic acids.

The synergistic herbicidal mixture particularly preferably comprises at least one herbicidal compound from amongst the group:

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quinclorac.

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cycloxydim, sethoxydim, clodinafop (and, if appropriate, cloquintocet), fenoxaprop-ethyl, fenoxaprop-P-ethyl, imazapyr, imazaquin, imazamethabenz, imazethapyr, pyrithiobac-sodium, metosulam, halosulfuron-methyl, nicosulfuron, N-[[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin-2-yl]amino]carbonyl]-2-(trifluoromethyl)-benzenesulfonamide, flufenacet, 2,4-D, diflufenzopyr, isoxaflutole, sulcotrione, glyphosate, glufosinate-ammonium, dimethenamid, S-metolachlor, benthiocarb, pendimethalin, acifluorfen, carfentrazone-ethyl, cinidon-ethyl, MCPA, dicamba and

- Also preferably, the synergistic herbicidal mixture

according to the invention comprises, as component B), at
least one herbicidal compound from amongst the group B12.

The synergistic herbicidal mixture according to the invention comprises in particular at least one herbicidal compound from amongst the group: propanil, pyridate, benzothiadiazinones, dinitrophenols, dipyridylenes, ureas, phenols, chloridazon, triazines, triazinones, uracils and biscarbamates.

Particularly preferably, the synergistic herbicidal mixture according to the invention comprises at least one herbicidal compound from amongst the group:

pyridate, bentazone, paraquat-dichloride, diuron, isoproturon, bromoxynil, chloridazon, atrazine or metribuzin.

Also particularly preferably, the herbicidal mixture according to the invention comprises at least one herbicidal compound from amongst the group: propanil, pyridate, dinitrophenols, dipyridylenes, chloridazon, triazinones, uracils and biscarbamates.

Particularly preferably, the synergistic herbicidal mixture according to the invention comprises at least one compound from amongst the group:

- pyridate, paraquat-dichloride, chloridazon or metribuzin.
- In a further particular embodiment, the synergistic herbicidal mixture according to the invention comprises, as component A), a 3-heterocyclyl-substituted benzoyl derivative of the formula I where
- is a heterocyclic radical selected from the group consisting of thiazol-2-yl, thiazol-4-yl and thiazol-5-yl, where the three abovementioned radicals may be unsubstituted or mono- or polysubstituted by halogen, C₁-C₄-alkyl, C₁-C₄-alkoxy, C₁-C₄-haloalkyl, C₁-C₄-haloalkoxy or C₁-C₄-alkylthio;
- and, as component B), at least one herbicidal compound from amongst the groups B1, B2, B4 to B12 or B14;
 - The synergistic herbicidal mixture according to the invention preferably comprises, as component B), at least one herbicidal compound from the following groups:
 - B1 acetyl-CoA carboxylase inhibitors (ACC):
 cyclohexenone oxime ethers or phenoxypropionic
 esters;
- B2 acetolactate synthase inhibitors (ALS):
 imidazolinones, pyrimidyl ethers, sulfonamides or
 sulfonylureas;
 - B4 auxin herbicides: pyridinecarboxylic acids or 2,4-D;
 - B5 auxin transport inhibitors;
 - B6 carotenoid biosynthesis inhibitors;

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- B7 enolpyruvylshikimate 3-phosphate synthase inhibitors;
- B8 glutamine synthetase inhibitors;

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- B9 lipid biosynthesis inhibitors: chloroacetanilides or thioureas,
- B10 mitosis inhibitors: dinitroanilines;
 - B11 protoporphyrinogen IX oxidase inhibitors: diphenyl ethers, oxadiazoles, cyclic imides or pyrazoles;

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- Photosynthesis inhibitors:

 pyridate, pyridafol, benzothiadiazinones,
 dipyridylenes, ureas, phenols, chloridazon,
 triazines or triazinones, in particular pyridate,
 benzothiadiazinones, dipyridylenes, ureas, phenols,
 chloridazon, triazines or triazinones;
- B14 growth substances:

 aryloxyalkanoic acids, benzoic acids or
 quinolinecarboxylic acids.

In particular, the synergistic herbicidal mixture according to the invention comprises, as component B), at least one herbicidal compound from the group:

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cycloxydim, sethoxydim, clodinafop (and, if appropriate, cloquintocet), fenoxaprop-ethyl, fenoxaprop-P-ethyl, imazapyr, imazaquin, imazamethabenz, imazethapyr, pyrithiobac-sodium, metosulam, halosulfuron-methyl, nicosulfuron, N-[[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin-2-yl]amino]carbonyl]-2-(trifluoromethyl)-benzenesulfonamide, flufenacet, 2,4-D, diflufenzopyr, isoxaflutole, sulcotrione, glyphosate, glufosinate-ammonium, dimethenamid, S-metolachlor, benthiocarb, pendimethalin, acifluorfen, carfentrazone-ethyl, cinidon-ethyl, pyridate, bentazon, paraquat-dichloride, diuron, isoproturon, bromoxynil, chloridazon, atrazine, metribuzin, MCPA, dicamba and

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quinclorac.

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5	Also preferably, the synergistic herbicidal mixture according to the invention comprises, as component B), at least one herbicidal compound from amongst the groups B1, B2, B4 to B11 and B14;
	In particular, the synergistic herbicidal mixture according to the invention comprises at least one herbicidal compound from amongst the following groups:
10	<pre>B1 acetyl-CoA carboxylase inhibitors (ACC): cyclohexenone oxime ethers or phenoxypropionic esters;</pre>
15	B2 acetolactate synthase inhibitors (ALS): imidazolinones, pyrimidyl ethers, sulfonamides or sulfonylureas;
20	B4 auxin herbicides: 2,4-D;
	B5 auxin transport inhibitors;
	B6 carotenoid biosynthesis inhibitors;
25	B7 enolpyruvylshikimate 3-phosphate synthase inhibitors;
	B8 glutamine synthetase inhibitors;
30	B9 lipid biosynthesis inhibitors: chloroacetanilides or thioureas,
35	B10 mitosis inhibitors: dinitroanilines;
	Bll protoporphyrinogen IX oxidase inhibitors: diphenyl ethers, oxadiazoles, cyclic imides or pyrazoles;
40	B14 growth substances: aryloxyalkanoic acid, benzoic acids or quinolinecarboxylic acids.
45	The synergistic herbicidal mixture particularly preferably comprises at least one herbicidal compound
	from amongst the group:

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cycloxydim, sethoxydim, clodinafop (and, if appropriate, cloquintocet), fenoxaprop-ethyl, fenoxaprop-P-ethyl, imazapyr, imazaquin, imazamethabenz, imazethapyr, pyrithiobac-sodium, metosulam, halosulfuron-methyl, nicosulfuron, N-{[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin-2-yl]amino]carbonyl]-2-(trifluoromethyl)-benzenesulfonamide, flufenacet, 2,4-D, diflufenzopyr, isoxaflutole, sulcotrione, glyphosate, glufosinate-ammonium, dimethenamid, S-metolachlor, benthiocarb, pendimethalin, acifluorfen, carfentrazone-ethyl, cinidon-ethyl, MCPA, dicamba and quinclorac.

- Also preferably, the synergistic herbicidal mixture
according to the invention comprises, as component B), at
least one herbicidal compound from amongst the group B12.

The synergistic herbicidal mixture according to the invention comprises in particular at least one herbicidal compound from amongst the group: propanil, pyridate, benzothiadiazinone, dinitrophenols, dipyridylenes, ureas, phenols, chloridazone, triazines, triazinones, uracils and biscarbamates.

Particularly preferably, the synergistic herbicidal mixture according to the invention comprises at least one herbicidal compound from amongst the group:

pyridate, bentazone, paraquat-dichloride, diuron, isoproturon, bromoxynil, chloridazon, atrazine or metribuzin.

Also particularly preferably, the herbicidal mixture according to the invention comprises at least one herbicidal compound from amongst the group: propanil, pyridate, dinitrophenols, dipyridylenes, chloridazon, triazinones, uracils and biscarbamates.

Particularly preferably, the synergistic herbicidal mixture according to the invention comprises at least one compound from amongst the group: pyridate, paraquat-dichloride, chloridazon or metribuzin.

In a further particular embodiment, the synergistic herbicidal mixture according to the invention comprises, as component A), a 3-heterocyclyl-substituted benzoyl derivative of the formula I where

R² is a heterocyclic radical selected from the group consisting of isoxazol-3-yl, isoxazol-4-yl and isoxazol-5-yl, where the three abovementioned radicals may be unsubstituted or mono- or polysubstituted by halogen, C₁-C₄-alkyl, C₁-C₄-alkoxy, C₁-C₄-haloalkyl, C₁-C₄-haloalkoxy or C₁-C₄-alkylthio;

and, as component B), at least one herbicidal compound from amongst the groups B1, B2, B4 to B12 or B14;

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The synergistic herbicidal mixture according to the invention preferably comprises, as component B), at least one herbicidal compound from the groups B1, B2, B4 to B11 and B14;

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In particular, the synergistic herbicidal mixture according to the invention comprises at least one herbicidal compound from the following groups:

- 20
- B1 acetyl-CoA carboxylase inhibitors (ACC):
 cyclohexenone oxime ethers or phenoxypropionic
 esters;
- B2 acetolactate synthase inhibitors (ALS):

 imidazolinones, pyrimidyl ethers, sulfonamides or
 sulfonylureas;
 - B4 auxin herbicides:
 pyridinecarboxylic acids or 2,4-D;

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- B5 auxin transport inhibitors;
- B6 carotenoid biosynthesis inhibitors;
- B7 enolpyruvylshikimate 3-phosphate synthase inhibitors;
 - B8 glutamine synthetase inhibitors;
- 40 B9 lipid biosynthesis inhibitors: chloroacetanilides or thioureas,
 - B10 mitosis inhibitors:
 dinitroanilines;

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Bll protoporphyrinogen IX oxidase inhibitors:

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diphenyl ethers, oxadiazoles, cyclic imides or pyrazoles;

B14 growth substances:
 aryloxyalkanoic acid, benzoic acids or
 quinolinecarboxylic acids.

Particularly preferably, the synergistic herbicidal mixture comprises at least one herbicidal compound from amongst the group:

cycloxydim, sethoxydim, clodinafop (and, if appropriate, cloquintocet), fenoxaprop-ethyl, fenoxaprop-P-ethyl, imazapyr, imazaquin, imazamethabenz, imazethapyr, pyrithiobac-sodium, metosulam, halosulfuron-methyl, nicosulfuron, N-[[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin-2-yl]amino]carbonyl]-2-(trifluoromethyl)-benzenesulfonamide, flufenacet, 2,4-D, diflufenzopyr, isoxaflutole, sulcotrione, glyphosate, glufosinate-ammonium, dimethenamid, S-metolachlor, benthiocarb, pendimethalin, acifluorfen, carfentrazone-ethyl, cinidon-ethyl, MCPA, dicamba and quinclorac.

Also preferably, the herbicidal mixture according to the invention comprises at least one herbicidal compound from amongst the group:

propanil, pyridate, dinitrophenols, dipyridylenes, chloridazon, triazinones, uracils and biscarbamates.

In particular, the synergistic herbicidal mixture according to the invention comprises at least one compound from amongst the group:

pyridate, paraquat-dichloride, chloridazon or metribuzin.

In a further particular embodiment, the synergistic herbicidal mixture comprises, as component A, a 3-heterocyclyl-substituted benzoyl derivative of the formula I and, as component B, a herbicidal compound. For particularly preferred embodiments, the preferences described above apply analogously.

In a further particular embodiment, the synergistic herbicidal mixture comprises, as component A, a
 3-heterocyclyl-substituted benzoyl derivative of the formula I and, as component B, two herbicidal compounds.

For particularly preferred embodiments, the preferences described above apply analogously.

- In a further particularly preferred embodiment, the synergistic herbicidal mixture comprises, as component B, a herbicidal compound, where with respect to the preferred embodiments the above preferences apply, and a herbicidal compound from amongst the groups B12 and B14.
- 10 The present invention also extends to herbicidal compositions which comprise a herbicidally active amount of a synergistic herbicidal mixture (comprising components A) and B) as described above), at least one liquid and/or solid carrier and, if desired, at least one surfactant.

The herbicidal compositions and synergistic herbicidal mixtures according to the invention can effect very good control of broad-leaved weeds and grass weeds in crops such as maize, cereals, rice and soya without damaging the crop plants, an 20 effect observed especially even at low rates of application.

Taking into consideration the variety of application methods in question, the herbicidal compositions and synergistic herbicidal mixtures according to the invention can additionally be employed

- 25 in a further number of crop plants for eliminating undesirable plants. Examples of suitable crops are the following:
 Allium cepa, Ananas comosus, Arachis hypogaea, Asparagus officinalis, Beta vulgaris spp. [sic] altissima, Beta vulgaris spp. [sic] rapa, Brassica napus var. napus, Brassica napus var.
- 30 napobrassica, Brassica rapa var. silvestris, Camellia sinensis, Carthamus tinctorius, Carya illinoinensis, Citrus limon, Citrus sinensis, Coffea arabica (Coffea canephora, Coffea liberica), Cucumis sativus, Cynodon dactylon, Daucus carota, Elaeis guineensis, Fragaria vesca, Glycine max, Gossypium hirsutum,
- 35 (Gossypium arboreum, Gossypium herbaceum, Gossypium vitifolium), Helianthus annuus, Hevea brasiliensis, Hordeum vulgare, Humulus lupulus, Ipomoea batatas, Juglans regia, Lens culinaris, Linum usitatissimum, Lycopersicon lycopersicum, Malus spp., Manihot esculenta, Medicago sativa, Musa spp., Nicotiana tabacum
- 40 (N.rustica), Olea europaea, Oryza sativa, Phaseolus lunatus, Phaseolus vulgaris, Picea abies, Pinus spp., Pisum sativum, Prunus avium, Prunus persica, Pyrus communis, Ribes sylvestre, Ricinus communis, Saccharum officinarum, Secale cereale, Solanum tuberosum, Sorghum bicolor (s. vulgare), Theobroma cacao,
- 45 Trifolium pratense, Triticum aestivum, Triticum durum, Vicia faba, Vitis vinifera und Zea mays.

Moreover, the herbicidal compositions and synergistic herbicidal mixtures according to the invention can also be used in crops which tolerate the action of herbicides due to breeding, including genetic engineering methods.

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The mixtures according to the invention, or the herbicidal compositions comprising them, can be employed, for example, in the form of directly sprayable aqueous solutions, powders, suspensions, also highly-concentrated aqueous, oily or other suspensions or dispersions, emulsions, oil dispersions, pastes, dusts, materials for spreading or granules, by means of spraying, atomizing, dusting, spreading or pouring.

The use forms depend on the intended purposes; in any case, they should guarantee the finest possible distribution of the active ingredients according to the invention.

Suitable inert auxiliaries are mineral oil fractions of medium to high boiling point such as kerosene and diesel oil, furthermore coal tar oils and oils of vegetable or animal origin, aliphatic, cyclic and aromatic hydrocarbons, e.g. paraffins, tetrahydronaphthalene, alkylated naphthalenes and their derivatives, alkylated benzenes and their derivatives, alcohols such as methanol, ethanol, propanol, butanol and cyclohexanol, ketones such as cyclohexanone, strongly polar solvents, such as N-methylpyrrolidone and water.

Aqueous use forms can be prepared from emulsion concentrates, suspensions, pastes, wettable powders or water-dispersible granules by adding water. To prepare emulsions, pastes or oil dispersions, the substrates [sic], as such or dissolved in an oil or solvent, can be homogenized in water by means of wetting agent, tackifier, dispersant or emulsifier. However, it is also possible to prepare concentrates composed of active substance, wetting agent, tackifier, dispersant or emulsifier and, if appropriate, solvent or oil, and these concentrates are suitable for dilution with water.

Suitable surfactants are the alkali metal, alkaline earth metal
40 and ammonium salts of aromatic sulfonic acids, e.g. ligno-,
phenol-, naphthalene- and dibutylnaphthalenesulfonic acid, and of
fatty acids, of alkyl- and alkylaryl sulfonates, of alkyl
sulfates, lauryl ether sulfates and fatty alcohol sulfates, and
salts of sulfated hexa-, hepta- and octadecanols, and of fatty
45 alcohol glycol ether, condensates of sulfonated naphthalene and
its derivatives with formaldehyde, condensates of naphthalene, or
of the naphthalenesulfonic acids, with phenol and formaldehyde,

polyoxyethylene octylphenyl ether, ethoxylated isooctyl-, octylor nonylphenol, alkylphenyl and tributylphenyl polyglycol ether,
alkylaryl polyether alcohols, isotridecyl alcohol, fatty
alcohol/ethylene oxide condensates, ethoxylated castor oil,
5 polyoxyethylene alkyl ethers or polyoxypropylene alkyl ethers,
lauryl alcohol polyglycol ether acetate, sorbitol esters,
lignin-sulfite waste liquors or methylcellulose.

Powders, materials for spreading and dusts can be prepared by 10 mixing or concomitantly grinding the synergistic herbicidal mixture or the individual active ingredients with a solid carrier.

- Granules, e.g. coated granules, impregnated granules and
 15 homogeneous granules, can be prepared by binding the active
 ingredients to solid carriers. Solid carriers are mineral earths
 such as silicas, silica gels, silicates, talc, kaolin, limestone,
 lime, chalk, bole, loess, clay, dolomite, diatomaceous earth,
 calcium sulfate, magnesium sulfate, magnesium oxide, ground
 20 synthetic material, fertilizers such as ammonium sulfate,
 ammonium phosphate, ammonium nitrate, ureas and products of
 vegetable origin such as cereal meal, tree bark meal, wood meal
 and nutshell meal, cellulose powders or other solid carriers.
- 25 The concentrations of the mixtures according to the invention in the ready-to-use products can be varied within wide ranges. In general, the formulations comprise from 0.01 to 95% by weight, preferably 0.5 to 90% by weight, of the mixture according to the invention.

The active ingredients of components A) and B) can be formulated jointly, but also separately, and/or applied to the plants, their environment and/or seeds jointly or separately. It is preferable to apply the active ingredients simultaneously. However, it is also possible to apply them separately.

Moreover, it may be advantageous to apply the herbicidal compositions and synergistic herbicidal mixtures according to the invention, jointly or separately, with additional other crop 40 protection agents, for example with pesticides or agents for controlling phytopathogenic fungi or bacteria. Also of interest is the miscibility with mineral salt solutions which are employed for treating nutritional and trace element deficiencies. Non-phytotoxic oils and oil concentrates can also be added.

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from the group of the aryloxyalkanoic acids in a weight ratio of 1:0.2 to 1:240, preferably 1:0,33 to 1:48.

Very particularly preferably, they comprise, as component B) fluoroxypyr in a weight ratio of 1:0.2 to 1:80, preferably 1:0.33 to 1:16.

Also very particularly preferably, they comprise, as component B), MCPA or mecoprop-P in a weight ratio of 1:1.6 to 1:240, preferably 1:2.67 to 1:48.

- Also especially preferably, the mixtures according to the invention comprise a 3-heterocycly1-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the benzoic acids, in particular dicamba, in a weight ratio of 1:0.3 to 1:160, preferably 1:0.5 to 1:32.
- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the quinolinecarboxylic acids, in particular quinclorac, in a weight ratio of 1:0,1 to 1:120, preferably 1:0.16 to 1:24.

Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group B16 (various other herbicides), in particular triaziflam, in a weight ratio of 1:0.2 to 1:150, preferably 1:0.3 to 1:30.

- Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and two herbicidal compounds from the groups B1 to B16, where the weight ratio of the 3-heterocyclyl-substituted benzoyl derivative of the formula I to each of the individual herbicidal components of B) is in the ranges described above.
- Especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I, a herbicidal compound from the group B2 and a herbicidal compound from the group B14 in a weight ratio of 1:0.004:0.1 to 1:160:240, preferably 1:0.006:0.16 to 1:32:48.

- Especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I, a herbicidal compound from the group B5 and a herbicidal compound from the group B14 in a weight ratio of 1:0.06:0.1 to 1:20:240, preferably 1:0.1:0.16 to 1:4:48.
- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I, a herbicidal compound from the group B9 and a herbicidal compound from the group B12 in a weight ratio of 1:0.24:0.12 to 1:80:800, preferably 1:0.48:0.2 to 1:16:160.
- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I, a herbicidal compound from the group B12 and a herbicidal compound likewise from the group B12 in a weight ratio of 1:0.12:0.12 to 1:800:800, preferably 1:0.2:0.2 to 1:160:160.
- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I, a herbicidal compound from the group B12 and a herbicidal compound from the group B14 in a weight ratio of 1:0.12:0.1 to 1:800:240, preferably 1:0.2:0.16 to 1:160:48.
- The rate of application of pure synergistic herbicidal mixture, 30 i.e. without formulation auxiliaries, amounts to 2 to 5000 g/ha, preferably 2 to 4500 g/ha, in particular 8 to 4500 g/ha, of active substance (a.s.), depending on the intended aim, the season, the target plants and growth stage.
- 35 The rate of application of 3-heterocyclyl-substituted benzoyl derivative of the formula I is 0.1 to 250 g/ha, as a rule 5 to 250 g/ha, preferably 25 to 150 g/ha, of active substance (a.s.).
- The preferred rate of application of the individual classes of 40 active ingredients, or of the active ingredients of component B, are compiled in Table 2.

Rate of application 100-400 100-400 100-400 100-400 25-400 25-300 25-100 100-800 50-300 25-150 (g/ha) 20-800 30-400 50-300 30-150 20-120 1-800 2-120 2-120 Active ingredient clodinafpop-P-propargyla fenoxaprop-P-ethyl pyrithiobac-sodium fenoxaprop-ethyl imazamethabenz imazaethopyr tralkoxydim cycloxydim sethoxydim imazaquin imazamox imazapyr phenoxyphenoxypropionic esters Class of active ingredient cyclohexenone oxime ethers pyrimidyl ethers imidazolinones Acetolactate synthase inhibitors (ALS) Acetyl-CoA carboxylase inhibitors Component B Bl **B**2

Table 2

Rate of application (g/ha)	1-225	1-20	25-225	1-60	1-120	5-120	1-120	10-120	10-120	5-120	10-60	10-60		5-120		10-60	250-2000	250-2000	25-750
Active ingredient		florasulam	flumetsulam	metosulam		halosulfuron-methyl	nicosulfuron	primisulfuron-methyl	prosulfuron	rimsulfuron	thifensulfuron-methyl	tribenuron-methyl	N-[[[4-methoxy-6-(trifluoromethyl)-	1,3,5-triazin-2-yl]amino]carbonyl]-	2-(trifluoromethyl)benzenesulfonamide	sulfosulfuron		fluthiamide	
Class of active ingredient	sulfonamides				sulfonylureas														
Component B																	Amides		Auxin herbicides
														•			B3		B4

Rate of application (g/ha)	25-750	25-750	50-750	15-100	15-100	25-600	25-200	100-600	25-300	25-200	25-300	360-1080	360-1080	360-1080	10-600	10-600	60-4000	60-4000
Active ingredient		clopyralid	2,4-D		diflufenzopyr		isoxaflutole	sulcotrione	mesotrione	isoxachlortole	ketospiradox		glyphosate	sulfosate		glufosinate-ammonium		
Class of active ingredient	pyridinecarboxylic acids						•	1			•					ı		chloroacetanilides
Component B				Auxin transport inhibitors		Carotenoid biosynthesis inhibitors						Enolpyruvylshikimat-3-phosphate synthase inhibitors (ESPS)			Glutamine synthetase inhibitors		Lipid biosynthesis inhibitors	
				BS		B6						B7			B8		B9	

			-) T								
Rate of application (g/ha)	60-2000	60-2000	250-4000	60-4000	60-4000	100-4000	1000-4000	375-3000	375-3000	375-3000	0.5-600	50-300	50-300	50-300	20-600	20-600	0.5-300	0.5-35
Active ingredient	dimethenamid	S-dimethenamid	acetochlor	metolachior	S-metolachlor		benthiocarb			pendimethalin			acifluorfen	acifluorfen-sodium		oxadiargyl		carfentrazone-ethyl
Class of active ingredient						thioureas			dinitroanilines			diphenyl ethers			oxadiazoles		cyclic imides	
Component B				·				Mitosis inhibitors			Protophorphyrinogen [sic] IX oxidase inhibitors							
								B10			B11							

									`) Z	,			_					
Rate of application (g/ha)	3-35	3-35	2-300	50-300	30-4000	250-1500	250-1000	480-1440	480-1440	100-800	. 100-800	250-1600	250-1600	250-1600	100-700	002-001	500-4000	250-4000	250-4000
Active ingredient	cinidon-ethyl	flumiclorac-pentyl	butafenacil	JV 485		pyridate	pyridafol		bentazone		paraquat-dichloride		diuron	isoprotoron		bromoxynił			atrazine
Class of active ingredient						•		benzothiadiazinones		dipyridylenes		ureas			phenois		chloridazon	triazines	
Component B					B12 Photosynthesis inhibitors														

·	_					-				3	·		_		-	
Rate of application (g/ha)	250-4000	30-300	30-300	500-1500	500-1500	500-1500	25-1200	50-1200	50-400	400-1200	400-1200	75-800	75-800	25-600	25-600	50-750
Active ingredient	terbutylazine		metribuzin			tridiphane			fluoroxypyr	MCPA	mecoprop-P		dicamba		quinclorac	triaziflam
Class of active ingredient		triazinone			oxiranes			aryloxyalkanoic acids				benzoic acids		quinolinecarboxylic acids		
Component B				Synergists			Growth substances									Various other herbicides
				B13		•	B14									B16

 $^{\mathrm{a}}$ If appropriate, 10-50 g/ha cloquintocet may also be added.

Use examples

- The mixtures according to the invention were applied pre- or post-emergence (foliar treatment). The herbicidal compounds of component B were applied in the formulation in which they are present as commercially available product.
- 10 Some of the experiments were greenhouse experiments and some were field trials on mini plots (on a site with sandy loam (pH 6.2 to 7.0) or sandy clay (pH 5.0 to 6.7) as the soil).
- The harmful plants differed with regard to size and developmental 15 state; on average, they were 5 to 20 cm long, depending on the growth habit.
- The herbicidally active compounds of components A) and B) were applied in succession or jointly, in the latter case in some cases as a tank mix and in some cases as a readymix, in the form of emulsions, aqueous solutions or suspensions, the vehicle being water (300 400 l/ha). In the case of the field trials, application was effected with the aid of a mobile plot sprayer.
- The test period extended over 3 to 8 weeks, and the stands were also observed at later points in time.
- Damage by the herbicidal compositions was evaluated with 30 reference to a scale of 0% to 100% in comparison with untreated control plots. 0 means no damage and 100 means complete destruction of the plants.
- The following examples will demonstrate the action of the 35 herbicidal compositions which can be used according to the invention, without excluding the possibility of other uses.
- In these examples, the value E at which only an additive effect of the individual active ingredients is to be expected was calculated by the method of S. R. Colby (Calculating synergistic and antagonistic responses of herbicide combinations, Weeds 15, 20 pp (1967).

This was done using the formula

$$E = X + Y - \frac{XY}{100}$$

where

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- X = Percentage of the herbicidal action of component A) at an
 application rate of a;
 - Y = Percentage of the herbicidal action of component B) at an application rate of b;

If the value observed exceeds the value E calculated in accordance with Colby's formula, then synergism is present.

The herbicidal mixtures according to the invention exert a greater herbicidal action than would have been expected according to Colby on the basis of the observed effects of the individual components when used alone.

The results of the tests are shown in Tables 3 to 82 below.

In these studies, the following plants were used:

30 Scientific name Common name Abutilon theophrasti Chinese lantern Alopecuros myosuroides Slender foxtail 35 Amaranthus retroflexus Redroot pigweed Anthemis mixta Camomile Bidens pilosa Common blackjack Brachiaria plantaginea Alexander grass Chenopodium album Lambsquarters 40 Cyperus iria Cyperus species Cyprus grass species Digitaria adscendens Crab grass Digitaria sanguinalis Hairy fingergrass Echinochloa crus-galli Common barnyard grass Galium aparine Bedstraw, catchweed Geranium carolinianum Carolina geranium

	Scientific name	Common name
	Ipomoea acuminata	Blue morning-glory
	Ipomoea lacunosa	-
5	Ipomoea purpurea var. diversifolia	-
	Ipomoea ssp. [sic]	Morning-glory species
	Lolium perenne	Perennial rye grass
	Panicum miliaceum	Prozo millet
	Phalaris spec.	Canary grass species
.0	Richardia brasiliensis	-
	Setaria faberi	Giant foxtail
	Setaria viridis	Green foxtail
	Sorghum bicolor	Common sorghum
5	Sorghum halepense	Johnson grass
	Stellaria media	Common chickweed
	Triticum aestivum	Winter wheat
	Veronica ssp. [sic]	Speedwell species
	Zea mays	Maize

Table 3: Herbicidal action of compound Ia.3 and "cycloxydim"

(B1) on Chenopodium album in the field (post-emergence treatment)

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		lication (g/ha	Damage (%)	Colby value E
	Ia.3	Cycloxydim		
	50		92	
30		100	0	
	50	100	98	92

Table 4: Herbicidal action of compound Ia.3 and "cycloxydim"

(B1) on Digitaria sanguinalis in the field

(post-emergence treatment)

		lication (g/has.)	Damage (%)	Colby value E
40	Ia.3	Cycloxydim	1	
	50		57	
		100	81	
	50	100	98	92

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Table 5: Herbicidal action of compound Ia.3 and "sethoxydim"
(B1) on Abutilon theophrasti in the field
(post-emergence treatment)

	• •	Damage (%)	Colby Value E
Ia.3	Sethoxydim	1	, , , , , ,
50		85	
	160	0	
50	160	94	85
	1a.3 50	50 160 50 160	a.s.) Damage (%) Ia.3 Sethoxydim 50 85 160 0 50 160 94

Table 6: Herbicidal action of compound Ia.3 and "sethoxydim"

(B1) on Setaria viridis in the field (post-emergence treatment)

	_	lication (g/ha.s.)	Damage (%)	Colby value E
20	Ia.3	Sethoxydim		
	100		75	
		160	93	
	100	160	. 99	98

Table 7: Herbicidal action of compound Ia.3 and "clodinafop-propargyl + cloquintocet" (B1) on Alopecurus myosuroides in the field (post-emergence treatment)

30				
		application ha a.s.)		
35	Ia.3	clodinafop- propargyl + cloquintocet	Damage (%)	Colby value E
33	7.5		10	## ## W
I		40	63	
	75	40	94	67

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Table 8: Herbicidal action of compound Ia.3 and "fenoxaprop-ethyl" (B1) on Alopecurus myosuroides in the field (post-emergence treatment)

5		application		
	Ia.3	fenoxaprop- ethyl	Damage (%)	Colby value E
	75		10	
10		83	82	P+-
	75	83	94	84

Table 9: Herbicidal action of compound Ia.3 and
"fenoxaprop-ethyl" (Bl) on Galium aparine in the field
(post-emergence treatment)

	Rate of application (g/ha a.s.)			
20	Ia.3	fenoxaprop- ethyl	Damage (%) Co	Colby value E
	75		63	
		83	0	
	75	83	75	63

Table 10: Herbicidal action of compound Ia.3 and "fenoxaprop-P-ethyl" (B1) on Amaranthus retroflexus in the greenhouse (post-emergence treatment)

30	Rate of application (g/ha a.s.)			
	Ia.3	fenoxaprop- ethyl	Damage (%)	Colby value E
	15.6		80	
		31.2	0	
35	15.6	31.2	95	80

Table 11: Herbicidal action of compound Ia.33 and "imazapyr" (B1) on Alopecurus myosuroides in the greenhouse (post-emergence treatment)

40	Rate of a	application a.s.)	Damage (%)	Colby value E
	Ia.33	imazapyr		
	15.6		40	
45		250	90	
	15.6	250	95	94

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Table 12: Herbicidal action of compound Ia.33 and "imazapyr" (B1) on Ipomoea ssp. [sic] in the greenhouse (post-emergence treatment)

5	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.33	imazapyr]	ours, varue s
	3.9		50	
		62.5	85	
10	3.9	62.5	95	93

Table 13: Herbicidal action of compound Ia.3 and "imazaquin" (B2) on Bidens pilosa in the field (post-emergence treatment)

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1+1	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	imazaquin]	
	75		30	
20		150	45	
	75	150	95	62

Table 14: Herbicidal action of compound Ia.3 and "imazamethabenz" (B2) on Stellaria media in the field (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	imazamethabenz		
30	75		91	
		525	0	
	75	525	99	91

Table 15: Herbicidal action of compound Ia.3 and "imazethapyr"

(B2) on Ipomoea acuminata in the field (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
40	Ia.3	imazethapyr		
	75		25	
	*	70	33	
	75	70	95	50

Table 16: Herbicidal action of compound Ia.3 and "imazethapyr" (B2) on Ipomoea purpurea var. diversifolia in the field (post-emergence treatment)

5	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	imazethapyr		sorpy varue E
	75		93	
		70	58	
10	. 75	70	99	97

Table 17: Herbicidal action of compound Ia.33 and "pyrithiobac-sodium" (B2) on Echinochloa crus-galli in the greenhouse (post-emergence treatment)

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	Rate of application (g/ha a.s.)			
	Ia.33	pyrithiobac- sodium	Damage (%)	Colby value E
20	1.9		55	+
		7.8	10	
	1.9	7.8	75	59

Table 18: Herbicidal action of compound Ia.33 and "metosulam"

(B2) on Veronica ssp. [sic] im the greenhouse

(post-emergence treatment)

30	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.33	metosulam	1 '''	
	62.5		20	
		1.9	40	
	62.5	1.9	75	52

35 Table 19: Herbicidal action of compound Ia.33 and "halosulfuron-methyl" (B2) on Alopecurus myosuroides in the greenhouse (post-emergence treatment)

40	Rate of application (g/ha a.s.)			
	Ia.33	halosulfuron- methyl	Damage (%)	Colby value E
	62.5		40	. ++-
		31.2	45	***
45	62.5	31.2	85	67

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Table 20: Herbicidal action of compound Ia.33 and "halosulfuron-methyl" (B2) on Amaranthus retroflexus in the greenhouse (post-emergence treatment)

5	Rate of application (g/ha a.s.)			
	Ia.33	halosulfuron- methyl	Damage (%)	Colby value E
	7.8		70	
10		7.8	80	
	7.8	7.8	98	94

Table 21: Herbicidal action of compound Ia.33 and "nicosulfuron" (B2) on Ipomoea lacunosa in the field (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.33	nicosulfuron	on 69	, , , , ,
20	75	~=~	69	
		35	39	
	75	35	90	81

Table 22: Herbicidal action of compound Ia.50 and "nicosulfuron"
25 (B2) on Amaranthus retroflexus in the greenhouse
(post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
30	Ia.50	nicosulfuron		7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7
	3.9		10	
		1.9	65	
	3.9	1.9	80	69

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Table 23: Herbicidal action of compound Ia.33 and
"N-[[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin2-yl]amino]carbonyl]-2-(trifluoromethyl)benzenesulfonamide" (B2) on Setaria faberi in the field
(post-emergence treatment)

	Rate of application (g/ha a.s.)			
10	Ia.33	N-[[[4-methoxy-6-(trifluoro-methyl)-1,3,5-triazin-2-yl]-amino]carbo-nyl]-2-(trifluoro-methyl)benzenesulfonamide	Damage (%)	Colby value E
	75		65	w to 40
		50	0	
	75	50	73	65

Table 24: Herbicidal action of compound Ia.3 and "2,4-D" (B4) on Abutilon theophrasti in the greenhouse (post-emergence treatment)

25	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	2,4-D		
	15.6		70	
30		62.5	40	
50	15.6	62.5	85	82

Table 25: Herbicidal action of compound Ia.3 and "2,4-D" (B4) on

Amaranthus retroflexus in the greenhouse (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby Value E
40	Ia.3	2,4-D		
-	15.6	* = -	55	
		62.5	20	
	15.6	62.5	70	64

Table 26: Herbicidal action of compound Ia.3 and "2,4-D" (B4) on Phalaris spec. in the field (post-emergence treatment)

5		application a.s.)	Damage (%)	Colby value E
	Ia.3	2,4-D		
	75		20	
		500	. 20	
	75	500	43	36

Table 27: Herbicidal action of compound Ia.3 and "isoxaflutole" (B6) on ipomoea ssp. [sic] in the greenhouse (post-emergence treatment)

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	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	isoxaflutole		
	31.2		75	
20		62.5	55	
	31.2	62.5	90	89

Table 28: Herbicidal action of compound Ia.3 and "isoxaflutole"

(B6) on Setaria viridis in the greenhouse
(post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
30	Ia.3	isoxaflutole		
	15.6		80	
ſ		31.2	. 30	
ſ	15.6	31.2	90	86

Table 29: Herbicidal action of compound Ia.3 and "sulcotrione"
(B6) on Ipomoea acuminata in the field (post-emergence treatment)

40		application a a.s.)	Damage (%)	Colby value E
	Ia.3	sulcotrione		
	75		25	
		300	86	
45	75	300	98	90

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Table 30: Herbicidal action of compound Ia.50 and "sulcotrione" (B6) on Amaranthus retroflexus in the greenhouse (post-emergence treatment)

5	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.50	sulcotrione		Corpy varue E
	31.2		60	
		250	45	
10	31.2	250	80	78

Table 31: Herbicidal action of compound Ia.3 and "glyphosate" (B7) on Geranium carolinianum in the field (post-emergence treatment)

		application a a.s.)	Damage (%)	Colby value E
	Ia.3	glyphosate	, , ,	dozzy varac z
0	150	***	30	
		840	97	
	150	840.	100	98

25 Table 32: Herbicidal action of compound Ia.33 and "glyphosate" (B7) on Sorghum halepense in the field (post-emergence treatment)

30	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.33	glyphosate	1	Tozz, varac L
	75		78	
		840	74	
	75	840	97	94
35		33.3 C 10.3 E 2. C 10.5 E 2		

Table 33: Herbicidal action of compound Ia.3 and

"glufosinate-ammonium" (B8) on Digitaria adscendens in the field (post-emergence treatment)

40	Rate of	application ha a.s.)		
	Ia.3	glufosinate- ammonium	Damage (%)	Colby value E
45	75		90	
. [400	75	
	75	400	100	98

Table 34: Herbicidal action of compound Ia.33 and "glufosinate-ammonium" (B8) on Echinochloa crus-galli in the greenhouse (post-emergence treatment)

5	Rate of application (g/ha a.s.)			
	Ia.33	glufosinate- ammonium	Damage (%)	Colby value E
	15.6		90	77-
-10		15.6	0	
	15.6	15.6	98	90

Table 35: Herbicidal action of compound Ia.3 and

"glufosinate-ammonium" (B8) on Ipomoea acuminata in the field (post-emergence treatment)

	Rate of application (g/ha a.s.)			
20	Ia.3	glufosinate- ammonium	Damage (%)	Colby value E
	75		25	
		400	75	
	75	400	98	81

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Table 36: Herbicidal action of compound Ia.33 and "glufosinate-ammonium" (B8) on Setaria faberi in the greenhouse (post-emergence treatment)

30	Rate of application (g/ha a.s.)			
	Ia.33	glufosinate- ammonium	Damage (%)	Colby value E
35	7.8		90	
35		31.2	65	
	7.8	31.2	98 .	96

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Table 37: Herbicidal action of compound Ia.3 and "flufenacet" (B3) on Digitaria adscendens in the field (post-emergence treatment)

5	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	flufenacet	J	7015, 14240 2
	75		90	
		600	58	
10	75	600	100	96

Table 38: Herbicidal action of compound Ia.3 and "dimethenamid" (B9) on Amaranthus retroflexus in the greenhouse (pre-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby-value E
	Ia.3	Dimethenamid		
20	31.2		40	
		125	80	
	31.2	125	100	88

25 Table 39: Herbicidal action of compound Ia.3 and "dimethenamid" (B9) on Cyperus iria in the greenhouse (pre-emergence treatment)

30	Rate of application (g/ha a.s.)		Damage (%)	Colby-value E
	Ia.3	Dimethenamid		
	31.2		50	
		62.5	95	
	31.2	62.5	100	98
35 h				والمراجع

Table 40: Herbicidal action of compound Ia.3 and "dimethenamid" (B9) on Digitaria sanguinalis in the greenhouse (pre-emergence treatment)

		plication (g/ha	Damage (%)	Colby value E
	Ia.3	Dimethenamid		
	62.5		60	
5		125	80	
	62.5	125	98	92

Table 41: Herbicidal action of compound Ia.33 and "dimethenamid" (B9) on Panicum miliaceum in the field (post-emergence treatment)

Rate of application (g/ha a.s.)		Damage (%)	Colby value E
Ia.33	dimethenamid	January (0)	COIDY VAIUE E
50		87	
	841	. 23	
50	841	94	90
	(g/i Ia.33 50	(g/ha a.s.) Ia.33 dimethenamid 50 841 50 841	(g/ha a.s.) Damage (%) Ia.33 dimethenamid 50 87 841 23 50 841 94

Table 42: Herbicidal action of compound Ia.33 and "dimethenamid" (B9) on Sorghum halepense in the field (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.33	dimethenamid		corpy varue E
20	75		78	
		1120	7	
	75	1120	90	80

25 Table 43: Herbicidal action of compound Ia.33 and "dimethenamid" (B9) on Veronica ssp. [sic] in the greenhouse (post-emergence treatment)

	Rate of application (g/ha a.s.)		Colby value E
3	Dimethenamid	Damage (%)	COLDI VALUE D
6		60	
	500	70	
6	500	90	88
	5	5 500	5 500 90

Table 44: Herbicidal action of compound Ia.52 and "dimethenamid" (B9) on Amaranthus retroflexus in the greenhouse (post-emergence treatment)

		Damage (%)	Colby value E
Ia.52	Dimethenamid		Total value L
62.5		75	
	500	10	
62.5	500	100	78
	Ia.52 62.5	62.5 500	a.s.) Damage (%) 1a.52 Dimethenamid 62.5 500 10

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Table 45: Herbicidal action of compound Ia.52 and "dimethenamid" (B9) on Veronica ssp. [sic] in the greenhouse (post-emergence treatment)

5		olication (g/ha	Damage (%)	Colby value E
	Ia.52	Dimethenamid		COIDY VAIUE E
	15.6		40	TO 100 AND 100
		500	70	
10	15.6	500	100	82

Table 46: Herbicidal action of compound Ia.33 and "acetochlor" (B9) on Abutilon theophrasti in the greenhouse (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.33	Ia.33 acetochlor	1	corpy varue E
20	7.8		90	
		31.2	0	
	7.8	31.2	100	90

25 Table 47: Herbicidal action of compound Ia.3 and "S-metolachlor" (B9) on Digitaria sanguinalis in the greenhouse (pre-emergence treatment)

Rate of application (g/ha a.s.)		Damage (%)	Colby value E
Ia.3	S-Metolachlor	j- (• /	Corpy varue E
62.5		60	***
	125	50	
62.5	125	85	80
	Ia.3 62.5	a.s.) Ia.3 S-Metolachlor 62.5 125 62.5 125	a.s.) Damage (%) Ia.3 S-Metolachlor 62.5 60 125 50 62.5 125 85

Table 48: Herbicidal action of compound Ia.3 and "S-metolachlor" (B9) on Echinochloa crus-galli in the greenhouse (pre-emergence treatment)

40		plication (g/ha a.s.)	Damage (%)	Colby value E
	Ia.3	S-Metolachlor		corpy value E
	62.5	~	60	
45		62.5	65	~
	62.5	62.5	98	86

Table 49: Herbicidal action of compound Ia.3 and "S-metolachlor" (B9) on Setaria viridis in the greenhouse (pre-emergence treatment)

5	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	S-Metolachlor	50 (0)	corpy value is
	15.6		20	
		62.5	70	
10	15.6	62.5	85	76

Table 50: Herbicidal action of compound Ia.33 and
"S-metolachlor" (B9) on Ipomoea ssp. [sic] in the
greenhouse (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.33	S-Metolachlor	go (• ,	Joseph value B
$^{\circ}$ Γ	62.5		80	
		62.5	. 0	
	62.5	62.5	90	80

25 Table 51: Herbicidal action of compound Ia.33 and "S-metolachlor" (B9) on Veronica ssp. [sic] in the greenhouse (post-emergence treatment)

30	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.33	S-Metolachlor		"""
	62.5		80	
	~~~	125	0	
35	62.5	125	98	80

Table 52: Herbicidal action of compound Ia.16 and "benthiocarb" (B9) on Cyperus iria in the field (post-emergence treatment)

		application	Damage (%)	Colby value E
	Ia.16	benthiocarb		10000
45	75		60	
		3000	50	
	75	3000	92	80

The mixtures according to the invention and the herbicidal compositions can be applied pre- or post-emergence. If the active ingredients are less well tolerated by certain crop plants, application techniques may be used in which the herbicidal 5 compositions are sprayed, with the aid of the spray apparatus, in such a way that they come into as little contact, if any, with the leaves of the sensitive crop plants while reaching the leaves of undesirable plants which grow underneath, or the bare soil (post-directed, lay-by).

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In the case of a post-emergence treatment of the plants, the herbicidal compositions according to the invention are preferably applied by foliar application. Application may be effected, for example, by usual spraying techniques with water as the carrier, 15 using amounts of spray mixture of approx. 100 to 1000 l/ha. The compositions may also be applied by the so-called "low-volume" and "ultra-low-volume" methods, or in the form of so-called granules.

20 As a rule, the synergistic herbicidal mixtures comprise components A) and B) in such weight ratios that the synergistic effect takes place. The ratios of component A) and B) in the mixture preferably range from 1:0.002 to 1:800, preferably from 1:0.003 to 1:160, particularly preferably from 1:0.02 to 1:160.

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- In particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the Group B1 (acetyl-CoA carboxylase inhibitors (ACC)) in a weight ratio of 1:0.1 to 1:80, preferably of 1:0.17 to 1:16.
- The mixtures according to the invention especially preferably comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the cyclohexenone oxime ethers, 35 preferably cycloxydim, sethoxydim or tralkoxydim, in particular sethoxydim or tralkoxydim, in a weight ratio of 1:0.4 to 1:80, preferably 1:0.67 to 1:16.
- 40 Also, the mixtures according to the invention especially preferably comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the phenoxyphenoxypropionic esters in a weight ratio of 1:0.1 to 1:60, preferably from 1:0.17 to 45 1:12.

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Very particularly preferably, they comprise, as component B), clodinafop-propargyl in a weight ratio of 1:0.1 to 1:20, preferably 1:0.17 to 1:4.

- Also very particularly preferably, they comprise, as component B), fenoxaprop-ethyl in a weight ratio of 1:0.2 to 1:60, preferably 1:0.34 to 1:12.
- Also very particularly preferably, they comprise, as component B), fenoxaprop-P-ethyl in a weight ratio of 1:0.1 to 1:30, preferably 1:0.16 to 1:6.
- Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the Group B2 (acetolactate synthase inhibitors) in a weight ratio of 1:0.004 to 1:160, preferably 1:0.006 to 1:32.
- Especially preferably, the mixtures according to the invention comprise 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the imidazolinones in a weight ratio of 1:0.08 to 1:160, preferably 1:0.13 to 1:32.
- Very particularly preferably, they comprise, as component B), imazapyr in a weight ratio of 1:0.12 to 1:80, preferably 1:0.2 to 1:16.
- Also very particularly preferably, they comprise, as component B), imazaquin in a weight ratio of 1:0.2 to 1:60, preferably 1:0.33 to 1:12.
  - Also very particularly preferably, they comprise, as component B), imazamethabenz in a weight ratio of 1:0.4 to 1:160, preferably 1:0.66 to 1:32.

Also very particularly preferably, they comprise, as component B), imazethapyr in a weight ratio of 1:0.12 to 1:30, preferably 1:0.2 to 1:6.

Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the pyrimidyl ethers, in particular pyrithiobac-sodium, in a weight ratio of 1:0.008 to 1:24, preferably 1:0.013 to 1:4.8.

- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from amongst the group of the sulfonamides in a weight ratio of 1:0.004 to 1:45, preferably 1:0.006 to 1:9.

Very particularly preferably, they comprise, as component B), flumetsulam in a weight ratio of 1:0.1 to 1:45, preferably 1:0.17 to 1:9.

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Also very particularly preferably, they comprise, as component B), metosulam in a weight ratio of 1:0.004 to 1:12, preferably 1:0.006 to 1:2.4.

- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the sulfonylureas in a weight ratio of 1:0.004 to 1:24, preferably 1:0.006 to 1:4.8.

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Very particularly preferably, they comprise, as component B), halosulfuron-methyl, rimsulfuron or N-[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin-2-yl]amino]carbonyl]-2-(trifluoromethyl)benzenesulfonamide in a weight ratio of 1:0.02 to 1:24, preferably 1:0.03 to 1:4.8.

Also very particularly preferably, they comprise, as component B), nicosulfuron in a weight ratio of 1:0.02 to 1:24, preferably 1:0.03 to 1:4.8.

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Also very particularly preferably, they comprise, as component B), primisulfuron-methyl or prosulfuron in a weight ratio of 1:0.04 to 1:24, preferably 1:0.06 to 1:4.8.

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Also very particularly preferably, they comprise, as component B), thisensulfuron-methyl, tribenuron-methyl or sulfosulfuron in a weight ratio of 1:0.04 to 1:12, preferably 1:0.06 to 1:2.4.

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Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the Group B3 (amides), in particular fluthiamide, in a weight ratio of 1:1 to 1:400, preferably 1:0.6 to 1:80.

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- Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the Group B4 (auxin-herbicides) in a weight ratio of 1:0.1 to 1:150, preferably 1:0.67 to 1:30.
- Especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from amongst the group of the pyridinecarboxylic acids, in particular clopyralid, in a weight ratio of 1:0.1 to 1:150, preferably 1:0.67 to 1:30.
- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and 2,4-D in a weight ratio of 1:0.2 to 1:150, preferably 1:0.33 to 1:30.
- Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the Group B5 (auxin transport inhibitors), preferably diflufenzopyr, in a weight ratio of 1:0.06 to 1:20, preferably 1:0.1 to 1:4.
- Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the Group B6 (carotenoid biosynthesis inhibitors) in a weight ratio of 1:0.1 to 1:120, preferably 1:0.17 to 1:24.
  - Especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and isoxaflutole or isoxachlortole in a weight ratio of 1:0.1 to 1:40, preferably 1:0.17 to 1:8.
- Also especially preferred, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and mesotrione or ketospiradox in a weight ratio of 1:0.1 to 1:60, preferably 1:0.16 to 1:12.
- Also especially preferred, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and sulcotrione in a weight ratio of 1:0.4 to 1:120, preferably 1:0.66 to 1:24.

- Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the Group B7 (enolpyruvylshikimate-3-phosphate synthase inhibitors
   (ESPS)), preferably glyphosate or sulfosate, in a weight ratio of 1:1.4 to 1:216, preferably 1:2.4 to 1:43.2.
- Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group B8 (glutamine synthetase inhibitors), preferably glufosinate-ammonium, in a weight ratio of 1:0.04 to 1:120, preferably 1:0.06 to 1:24.
- 15 Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group B9 (lipid biosynthesis inhibitors) in a weight ratio of 1:0.24 to 1:800, preferably 1:0.40 to 1:160.
- Especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the chloroacetanilides in a weight ratio of 1:0.24 to 1:800, preferably 1:0.4 to 1:160.
- Very particularly preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and dimethenamid or S-dimethenamid in a weight ratio of 1:0.24 to 1:400, preferably 1:0,4 to 1:80.
- Also very particularly preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and acetochlor in a weight ratio of 1:1 to 1:800, preferably 1:1.67 to 1:160.
- Also very particularly preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and metolachlor or S-metolachlor in a weight ratio of 1:0.24 to 1:800, preferably 1:0.40 to 1:160.
- Especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound

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from the group of the thioureas in a weight ratio of 1:0.4 to 1:800, preferably 1:0.66 to 1:160.

Very particularly preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and benthiocarb in a weight ratio of 1:4 to 1:800, preferably 1:6.6 to 1:160.

- * Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group B10 (mitosis inhibitors), preferably a dinitroaniline, in particular pendimethalin, in a weight ratio of 1:1,5 to 1:600, preferably 1:2,5 to 1:120.
- Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group Bl1 (protoporphyrinogen IX oxidase inhibitors) in a weight ratio of 1:0.002 to 1:120, preferably 1:0.003 to 1:24.
  - Especially preferably, the mixtures according to the invention comprise 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the diphenylethers, in particular acifluorfen or acifluorfen-sodium, in a weight ratio of 1:0.2 to 1:60, preferably 1:0.33 to 1:12.
- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the oxadiazoles, in particular oxadiargyl, in a weight ratio of 1:0.2 to 1:120, preferably 1:0.33 to 1:24.
  - Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the cyclic imides in a weight ratio of 1:0.002 to 1:60, preferably 1:0.003 to 1:12.

Very particularly preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and carfentrazone-ethyl in a weight ratio of 1:0.002 to 1:7, preferably 1:0.003 to 1:1.4.

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- Also very particularly preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I cinidon-ethyl or flumiclorac-pentyl, in a weight ratio of 1:0.012 to 1:7, preferably 1:0.02 to 1:1.4.

Also very particularly preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and butafenacil in a weight ratio of 1:0.02 to 1:60, preferably 1:0.03 to 1:12.

Also very particularly preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and JV 485 in a weight ratio of 1:0.2 to 1:60, preferably 1:0.3 to 1:12.

- In particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from group B12 (photosynthesis inhibitors) in a weight ratio of 1:0.12 to 1:800, preferably 1:0.2 to 1:160.
- Especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and pyridate or pyridafol in a weight ratio of 1:1 to 1:300, preferably 1:1.67 to 1:60.
- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the benzothiadiazinones, in particular bentazone, in a weight ratio of 1:1.92 to 1:288, preferably 1:3.2 to 1:57.6.
- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the dipyridylenes, in particular paraquat-dichloride, in a weight ratio of 1:0.4 to 1:160, preferably 1:0.66 to 1:32.
- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the ureas, in particular diuron or

isoproturon, in a weight ratio of 1:1 to 1:320, preferably 1:1.67 to 1:64.

- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the phenols, in particular bromoxynil, in a weight ratio of 1:0.4 to 1:140, preferably 1:0.67 to 1:28.

Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and chloridazon in a weight ratio of 1:2 to 1:800, preferably 1:3.3 to 1:160.

- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the triazines, in particular atrazine or terbutylazine, in a weight ratio of 1:1 to 1:800, preferably 1:1.67 to 1:160.
- Also especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group of the triazinones, in particular metribuzin, in a weight ratio of 1:0.12 to 1:60, preferably 1:0.2 to 1:12.
- 30 Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group B13 (synergists), preferably an oxirane, in particular tridiphane, in a weight ratio of 1:2 to 1:300, preferably 1:3.33 to 1:60.
- Also in particular, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound from the group B14 (growth substances) in a weight ratio of 1:0.1 to 1:240, preferably 1:0.167 to 1:48.
- Especially preferably, the mixtures according to the invention comprise a 3-heterocyclyl-substituted benzoyl derivative of the formula I and a herbicidal compound

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Table 53: Herbicidal action of compound Ia.3 and "pendimethalin" (Bl0) on Brachiaria plantaginea in the field (post-emergence treatment)

5	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	pendimethalin		corpy varue E
	75	***	96	
		990	0	
10	75	990	98	96

Table 54: Herbicidal action of compound Ia.3 and "acifluorfen" (B11) on Galium aparine in the field (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	acifluorfen		corpy value E
20	75		60	
		100	48	
	75	100	95	79

25 Table 55: Herbicidal action of compound Ia.33 and "carfentrazone-ethyl" (Bl1) on Amaranthus retroflexus in the greenhouse (post-emergence treatment)

30	Rate of application (g/ha a.s.)			
	Ia.33	carfentrazone- ethyl	Damage (%)	Colby value E
	1.9		30	
		0.9	60	
35	1.9	0.9	90	72

Table 56: Herbicidal action of compound Ia.3 and "carfentrazone-ethyl" (Bl1) on Anthemis mixta in the field (post-emergence treatment)

	Rate of application (g/ha a.s.)			
4.5	Ia.3	carfentrazone- ethyl	Damage (%)	Colby value E
15	75		68	W W W
		30	0	
	75	. 30	91	68

Table 57: Herbicidal action of compound Ia.33 and "cinidon-ethyl" (B11) on Galium aparine in the field (post-emergence treatment)

5	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.33	cinidon-ethyl	Jamage (1)	COIDY Value E
	1.9		20	***
		7.8	90	
10	1.9	7.8	100	92

Table 58: Herbicidal action of compound Ia.3 and "pyridate" (B12) on Bidens pilosa in the field (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	pyridate	s- (o,	corpy value E
20	75		25	
		450	25	
	75	450	96	44

25 Table 59: Herbicidal action of Ia.3 and "pyridate" (B12) on Setaria faberi in the field (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
30	Ia.3	pyridate		colly value L
	75		99	
		450	0	
	75	450	100	99

Table 60: Herbicidal action of compound Ia.3 and "bentazone" (B12) on Richardia brasiliensis in the field (post-emergence treatment)

Rate of application (g/ha a.s.)			Colby value E
Ia.3	Bentazone	1	Joseph value E
75	**	70	
	1440	77	
. 75	1440	99	93
	Rate of app a Ia.3	Rate of application (g/ha a.s.)  Ia.3 Bentazone  75 1440  75 1440	Rate of application (g/ha a.s.)  Damage (%)  1a.3  Bentazone  75   1440  77  75  1440  99

Table 61: Herbicidal action of compound Ia.3 and "paraquat-dichloride" (B12) on Lolium perenne in the field (post-emergence treatment)

5	Rate of application (g/ha a.s.)		_	
	Ia.3	paraquat- dichloride	Damage (%)	Colby value E
	75		10	
0		400	97	
	75	400	100	97

Table 62: Herbicidal action of compound Ia.33 and "diuron" (B12)

on Alopecurus myosuroides in the greenhouse
(post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
20	Ia.33	diuron		coiby value E
	62.5		40	100 000 000
		250 ·	80	
	62.5	250	95	88

Table 63: Herbicidal action of compound Ia.3 and "isoproturon"

(B12) on Stellaria media in the field (post-emergence treatment)

30	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	isoproturon		corpl varde F
	75		91	
		1000	94	
35	75	1000	100	99

Table 64: Herbicidal action of compound Ia.3 and "bromoxynil" (B12) on Galium aparine in the field (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	bromoxynil		corpy value E
5	75		60	
		470	84	
	75	470	98	94

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Table 65: Herbicidal action of compound Ia.3 and "chloridazon" (B12) on Ipomoea purpurea var. diversifolia in the field (post-emergence treatment)

5	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	chloridazon		corby value E
	75		94	
		1720	40	
10	75	1720	100	96

Table 66: Herbicidal action of compound Ia.3 and "atrazine" (B12) on Abutilon theophrasti in the field (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	atrazine	]	Colby value E
20	75		85	
		1120	32	
	75	1120	96	90

25 Table 67: Herbicidal action of compound Ia.3 and "atrazine" (B12) on Setaria faberi in the field (post-emergence treatment)

30	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	atrazine	]	COLDY VALUE E
	75	***	95	
		1120	20	
	75	1120	99	96

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Table 68: Herbicidal action of compound Ia.33 and "atrazine" (Bl2) on Sorghum bicolor in the field (post-emergence treatment)

40		application a a.s.)	Damage (%)	Colby value E
	Ia.33	atrazine	]	Joseph Agrae E
	· 75		78	***
45		840	27	
L	75	840	90	84

Table 69: Herbicidal action of compound Ia.3 and "metribuzin" (Bl2) on Bidens pilosa in the field (post-emergence treatment)

5	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	metribuzin	1	corpl varue F
	75		25	***
		200	38	
10	75	200	73	54

Table 70: Herbicidal action of compound Ia.3 and "metribuzin"

(B12) on Cyperus species in the field (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	metribuzin	, , , , , , , , , , , , , , , , , , , ,	corpy varue E
0	75		5	
		200	50	
	75	200	75	53

25 Table 71: Herbicidal action of compound Ia.3 and "MCPA" (B14) on Cyperus species in the field (post-emergence treatment)

30	Rate of application (g/ha a.s.)		Damage (%)	Colby value E
	Ia.3	MCPA.		colly value in
	75		0	
		600	5	
	75	600	48	5

Table 72: Herbicidal action of compound Ia.16 and "dicamba" (B14) on Amaranthus retroflexus in the field (post-emergence treatment)

40	Rate of a (g/ha	pplication a.s.)	Damage (%)	Colby value E
	Ia.16	dicamba	7 '''	1 10200
	100	** **	96	
45		280	25	
	100	280	100	97

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Table 73: Herbicidal action of compound Ia.33 and "dicamba" (B14) on Sorghum bicolor in the field (post-emergence treatment)

Rate of a (g/ha	pplication a.s.)	Damage (%)	Colhum
Ia.33	dicamba		Colby value E
75	- m 4-	78	
	560	17	
75	560	89	01
	(g/ha Ia.33 75	75 560	(g/ha a.s.)     Damage (%)       Ia.33     dicamba       75      78        560     17

Table 74: Herbicidal action of compound Ia.3 and "quinclorac" (B14) on Ipomoea ssp. [sic] in the greenhouse (post-emergence treatment)

	Rate of application (g/ha a.s.)		Damage (%)	Colbu walus 7
	Ia.3	Quinclorac	1 544.1290 (1)	Colby value E
	31.2		75	
0		250	70	
	31.2	250	100	93

Table 75: Herbicidal action of compound Ia.3 and "quinclorac"

(B14) on Veronica ssp. [sic] in the greenhouse (post-emergence treatment)

	lication (g/ha .s.)	Damage (%)	Colby walve 5
Ia.3	Quinclorac	Jamage (0)	Colby value E
31.2		80	
***	500	80	
31.2	500	100	96
	Ia.3 31.2	Ia.3 Quinclorac 31.2 500 31.2 500	Ia.3     Quinclorac       31.2      80        500     80       31.2     500     100

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Table 76: Herbicidal action of compound Ia.3, "nicosulfuron" (B2) and "dicamba" (B14) on Ipomoea acuminata in the field (post-emergence treatment)

40		application a a.s.)		
	Ia.3	nicosulfuron + dicamba	Damage (%)	Colby value E
į	75		23	
45		20 + 192	89	
	75	20 + 192	97	92

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Table 77: Herbicidal action of compound Ia.3, "diflufenzopyr" (B5) and "dicamba" (B14) on Echinochloa crus-galli in the field (post-emergence treatment)

5	Rate of application (g/ha a.s.)			
:	Ia.3	diflufenzopyr + dicamba	Damage (%)	Colby value E
	75		98 .	
10		56 + 140	5	
	75	56 + 140	99	98

Table 78: Herbicidal action of compound Ia.33, "diflufenzopyr"

(B5) and "dicamba" (B14) on Sorghum halepense in the field (post-emergence treatment)

	Rate of application (g/ha a.s.)			
20	Ia.33	diflufenzopyr + dicamba	Damage (%)	Colby value E
	75		78	
		.60 + 150	27	
	75	60 + 150	90	84

Table 79: Herbicidal action of compound Ia.33, "dimethenamide" (B9) and "atrazine" (B12) on Sorghum halepense in the field (post-emergence treatment)

30	Rate of application (g/ha a.s.)			
	Ia.33	dimethenamide + atrazine	Damage (%)	Colby value E
, [	75		78	
35	***	840 + 960	5	
	7.5	840 + 960	97	79

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Table 80: Herbicidal action of compound Ia.3, "bentazone" (B12) and "atrazine" (B12) on Brachiaria plantaginea in the field (post-emergence treatment)

5	Rate of application (g/ha a.s.)			
	Ia.3	bentazone + atrazine	Damage (%)	Colby value E
	75		95	
10	~ ~ ~	800 + 800	25	
	75	800 + 800	98	96

Table 81: Herbicidal action of compound Ia.33, "atrazine" (B12)

and "dicamba" (B14) on Ipomoea lacunosa in the field
(post-emergence treatment)

	Rate of application (g/ha a.s.)			
20	Ia.33	atrazine + dicamba	Damage (%)	Colby value E
	75		69	- w
		920 + 480	83	
	75	920 + 480	99	95

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Table 82: Herbicidal action of compound Ia.33, "atrazine" (B12) and "dicamba" (B12) on Setaria faberi in the field (post-emergence treatment)

30	Rate of application (g/ha a.s.)			
	Ia.33	atrazine + dicamba	Damage (%)	Colby value E
<b></b> [	75		65	
35		367 + 193	20	****
	75	367 + 193	89	72

Further experiments demonstrated that the mixtures according to 40 the invention are crop plant selective (Tables 83 and 84).

Table 83: Phytotoxicity of compound Ia.52 and "dimethenamid" (B9) to Triticum aestivum in the greenhouse (post-emergence treatment)

ا	Rate of application (g/ha a.s.)		Phytotoxicity (%)
	Ia.52	Dimethenamide	
	62.5		0
L		500	0
10	62.5	500	0

Table 84: Phytotoxicity of compound Ia.33 and "S-metolachlor"
(B9) on Zea mays in the greenhouse (post-emergence treatment)

		olication (g/ha	Phytotoxicity (%)
C	Ia.33	S-Metolachlor	
20	62.5	***	0
		125	. 0
	62.5	125	0

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We claim:

A synergistic herbicidal mixture comprising

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at least one 3-heterocyclyl-substituted benzoyl A) derivative of the formula I

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15 in which the variables have the following meanings:

> $R^1$ ,  $R^3$ are hydrogen, halogen, C1-C6-alkyl,  $C_1-C_6$ -haloalkyl,  $C_1-C_6$ -alkoxy,  $C_1-C_6$ -haloalkoxy, C₁-C₆-alkylthio, C₁-C₆-alkylsulfinyl or C₁-C₆-alkylsulfonyl;

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 $\mathbb{R}^2$ is a heterocyclic radical selected from the group: thiazol-2-yl, thiazol-4-yl, thiazol-5-yl, isoxazol-3-yl, isoxazol-4-yl, isoxazol-5-yl, 4,5-dihydroisoxazol-3-yl, 4,5-dihydroisoxazol-4-yl and

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4,5-dihydroisoxazol-5-yl, it being possible for the nine radicals mentioned to be unsubstituted or mono- or polysubstituted by halogen,

30

 $C_1-C_4-alkyl$ ,  $C_1-C_4-alkoxy$ ,  $C_1-C_4-haloalkyl$ , C₁-C₄-haloalkoxy or C₁-C₄-alkylthio;

R⁴

is hydrogen, halogen or C1-C6-alkyl;

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R⁵ is C₁-C₆-alkyl;

R6

is hydrogen or C₁-C₆-alkyl;

or one of its environmentally compatible salts;

40

and

a synergistically effective amount of at least one herbicidal compound from the group of the acetyl-CoA carboxylase inhibitors (ACC), acetolactate synthase 45 inhibitors (ALS), amides, auxin herbicides, auxin transport inhibitors, carotenoid biosynthesis inhibitors,

enolpyruvylshikimate 3-phosphate synthase inhibitors (ESPS), glutamine synthetase inhibitors, lipid biosynthesis inhibitors, mitosis inhibitors, protoporphyrinogen IX oxidase inhibitors, photosynthesis inhibitors, synergists, growth substances, cell wall biosynthesis inhibitors and a variety of other herbicides.

- A synergistic herbicidal mixture as claimed in claim 1
   comprising, as component B), at least one herbicidal compound from the groups B1 to B16:
  - Bl acetyl-CoA carboxylase inhibitors (ACC): cyclohexenone oxime ethers, phenoxyphenoxypropionic esters or arylaminopropionic acids;
  - B2 acetolactate synthase inhibitors (ALS): imidazolinones, pyrimidyl ethers, sulfonamides or sulfonylureas;

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- B3 amides;
- B4 auxin herbicides:

  pyridinecarboxylic acids, 2,4-D or benazolin;

- B5 auxin transport inhibitors;
- B6 carotenoid biosynthesis inhibitors;
- 30 B7 enolpyruvylshikimate 3-phosphate synthase inhibitors (ESPS);
  - B8 glutamine synthetase inhibitors;
- 35 B9 lipid biosynthesis inhibitors: anilides, chloroacetanilides, thioureas, benfuresate or perfluidone;
- B10 mitosis inhibitors:

  carbamates, dinitroanilines, pyridines, butamifos, chlorthal-dimethyl (DCPA) or maleic hydrazide;
  - B11 protoporphyrinogen IX oxidase inhibitors: diphenyl ethers, oxadiazoles, cyclic imides or pyrazoles;
- B12 photosynthesis inhibitors:

propanil, pyridate, pyridafol, benzothiadiazinones, dinitrophenols, dipyridylenes, ureas, phenols, chloridazon, triazines, triazinones, uracils or biscarbamates;

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- B13 synergists:
   oxiranes;
- B14 growth substances: aryloxyalkanoic acids, benzoic acids or quinolinecarboxylic acids;
  - B15 cell wall synthesis inhibitors:
- 15 B16 various other herbicides: dichloropropionic acids, dihydrobenzofurans, phenylacetic acids or aziprotryn, barban, bensulide, benzthiazuron, benzofluor, buminafos, buthidazole, buturon, cafenstrole, chlorbufam, xhlorofenprop-methyl, chloroxuron, 20 cinmethylin, cumyluron, cycluron, cyprazine, cyprazole, dibenzyluron, dipropetryn, dymron, eglinazin-ethyl, endothall, ethiozin, flucabazone, fluorbentranil, flupoxam, isocarbamid, isopropalin, karbutilate, mefluidide, monuron, napropamide, napropanilide, 25 nitralin, oxaciclomefone, phenisopham, piperophos, procyazine, profluralin, pyributicarb, secbumeton, sulfallate (CDEC), terbucarb, triazofenamide, triaziflam or trimeturon;
- or their environmentally compatible salts.
  - 3. A synergistic herbicidal mixture as claimed in claim 1 or 2, comprising, as component B), at least one herbicidal compound from the groups B1 to B16:

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- Bl acetyl-CoA carboxylase inhibitors (ACC):
  - cyclohexenone oxime ethers: alloxydim, clethodim, cloproxydim, cycloxydim, sethoxydim, tralkoxydim, butroxydim, clefoxydim or tepraloxydim;
  - phenoxyphenoxypropionic esters: clodinafop-propargyl (and, if appropriate, cloquintocet), cyhalofop-butyl, diclofop-methyl, fenoxaprop-ethyl, fenoxaprop-P-ethyl, fenthiapropethyl, fluazifop-butyl, fluazifop-P-butyl, haloxyfop-ethoxyethyl, haloxyfop-methyl,

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haloxyfop-P-methyl, isoxapyrifop, propaquizafop, quizalofop-ethyl, quizalofop-P-ethyl or quizalofop-tefuryl; or

- arylaminopropionic acids:

flamprop-methyl or flamprop-isopropyl;

### B2 acetolactate synthase inhibitors (ALS):

- imidazolinones: imazapyr, imazaquin, imazamethabenz-methyl (imazame), imazamoc, imazapic, imazethapyr or imazamethapyr;

pyrimidyl ethers: pyrithiobac-acid, pyrithiobac-sodium, bispyribacsodium, KIH-6127 or pyribenzoxym;

sulfonamides:

florasulam, flumetsulam or metosulam; or

sulfosulfuron or idosulfuron;

- sulfonylureas:
amidosulfuron, azimsulfuron, bensulfuron-methyl,
chlorimuron-ethyl, chlorsulfuron, cinosulfuron,
cyclosulfamuron, ethametsulfuron-methyl,
ethoxysulfuron, flazasulfuron, halosulfuron-methyl,
imazosulfuron, metsulfuron-methyl, nicosulfuron,
primisulfuron-methyl, prosulfuron,
pyrazosulfuron-ethyl, rimsulfuron,
sulfometuron-methyl, thifensulfuron-methyl,
triasulfuron, tribenuron-methyl,
triflusulfuron-methyl, N-[[[4-methoxy-6(trifluoromethyl)-1,3,5-triazin-2-yl]amino]carbonyl]-2-(trifluoromethyl)-benzenesulfonamide,

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#### B3 amides:

 allidochlor (CDAA), benzoylprop-ethyl, bromobutide, chlorthiamid, diphenamid, etobenzanid (benzchlomet), fluthiamide, fosamin or monalide;

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### B4 auxin herbicides:

- pyridine carboxylic acids: clopyralid or picloram; or
- 2,4-D or benazolin;

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# B5 auxin transport inhibitors:

naptalame or diflufenzopyr;

#### B6 carotenoid biosynthesis inhibitors:

45 - benzofenap, clomazone (dimethazone), diflufenican, fluorochloridone, fluridone, pyrazolynate, pyrazoxyfen, isoxaflutole, isoxachlortole,

mesotrione, sulcotrione (chlormesulone), ketospiradox, flurtamone, norflurazon or amitrol;

- B7 enolpyruvylshikimate-3-phosphate synthase inhibitors (ESPS):
  - glyphosate or sulfosate;
  - B8 Glutamine synthetase inhibitors:
    - bilanafos (bialaphos) or glufosinate-ammonium;

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Lipid biosynthesis inhibitors:

- anilides:
  - anilofos or mefenacet;
- chloroacetanilides:
  dimethenamid, S-dimethenamid, acetochlor, alachlor,
  butachlor, butenachlor, diethatyl-ethyl,
  dimethachlor, metazachlor, metolachlor,
  S-metolachlor, pretilachlor, propachlor, prynachlor,
  terbuchlor, thenylchlor or xylachlor;
- thioureas:
   butylate, cycloate, di-allate, dimepiperate, EPTC,
   esprocarb, molinate, pebulate, prosulfocarb,
   thiobencarb (benthiocarb), tri-allate or vernolate;
   or
- 25 benfuresate or perfluidone;

### B10 mitosis inhibitors:

- carbamates:
  - asulam, carbetamid, chlorpropham, orbencarb, pronamid (propyzamid), propham or tiocarbazil;
- dinitroanilines: benefin, butralin, dinitramin, ethalfluralin, fluchloralin, oryzalin, pendimethalin, prodiamine or trifluralin;
- pyridines:
   dithiopyr or thiazopyr; or
  - butamifos, chlorthal-dimethyl (DCPA) or maleic hydrazide;
- 40 Bll protoporphyrinogen IX oxidase inhibitors:
  - diphenyl ethers: acifluorfen, acifluorfen-sodium, aclonifen, bifenox, chlornitrofen (CNP), ethoxyfen, fluorodifen, fluoroglycofen-ethyl, fomesafen, furyloxyfen, lactofen, nitrofen, nitrofluorfen or oxyfluorfen;
  - oxadiazoles:
     oxadiargyl or oxadiazon;

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5		cyclic imides: azafenidin, butafenacil, carfentrazone-ethyl, cinidon-ethyl, flumiclorac-pentyl, flumioxazin, flumipropyn, flupropacil, fluthiacet-methyl, sulfentrazone or thidiazimin; or pyrazoles: ET-751, JV 485 or nipyraclofen;
	B12	photosynthesis inhibitors:
10		- propanil, pyridate or pyridafol;
		- benzothiadiazinones:
		bentazone;
		- dinitrophenols:
15		<pre>bromofenoxim, dinoseb, dinoseb-acetate, dinoterb or DNOC;</pre>
	•	- dipyridylenes:
		<pre>cyperquat-chloride, difenzoquat-methylsulfate, diquat or paraquat-dichloride;</pre>
		- ureas:
20		chlorbromuron, chlorotoluron, difenoxuron, dimefuron,
		diuron, ethidimuron, fenuron, fluometuron,
	•	isoproturon, isouron, linuron, methabenzthiazuron,
		methazole, metobenzuron, metoxuron, monolinuron,
		neburon, siduron or tebuthiuron;
25		- phenois:
		bromoxynil or ioxynil;
		- chloridazon;
		- triazines:
		ametryn, atrazine, cyanazine, desmetryn,
30		dimethamethryn, hexazinone, prometon, prometryn,
		propazine, simazine, simetryn, terbumeton, terbutryn,
		terbutylazine or trietazine;
		- triazinones:
	•	metamitron or metribuzine;
35		- uracils:
		bromacil, lenacil or terbacil; or
		- biscarbamates:
		desmedipham or phenmedipham;
40	B13	synergists:
		- oxiranes:
		tridiphane;
	B14	growth substances:

- aryloxyalkanoic acids:

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2,4-DB, clomeprop, dichlorprop, dichlorprop-P (2,4-DP-P), fluoroxypyr, MCPA, MCPB, mecoprop, mecoprop-P, or triclopyr;

- benzoic acids:
  - chloramben or dicamba; or
- quinolinecarboxylic acids: quinclorac or quinmerac;

# B15 cell wall synthesis inhibitors:

- isoxaben or dichlobenil;

## B16 various other herbicides

- dichloropropionic acids: dalapon;
- dihydrobenzofurans:
   ethofumesate;
  - phenylacetic acids:
     chlorfenac (fenac); or
- aziprotryn, barban, bensulide, benzthiazuron,
  benzofluor, buminafos, buthidazole, buturon,
  cafenstrole, chlorbufam, chlorfenprop-methyl,
  chloroxuron, cinmethylin, cumyluron, cycluron,
  cyprazine, cyprazole, dibenzyluron, dipropetryn,
  dymron, eglinazin-ethyl, endothall, ethiozin,
  flucabazone, fluorbentranil, flupoxam, isocarbamid,
  isopropalin, karbutilate, mefluidide, monuron,
  napropamide, napropanilide, nitralin, oxaciclomefone,
  phenisopham, piperophos, procyazine, profluralin,
  pyributicarb, secbumeton, sulfallate (CDEC),
  terbucarb, triazofenamid, triaziflan or trimeturon;

or their environmentally compatible salts.

- A synergistic herbicidal mixture as claimed in any of
   claims 1 to 3, comprising, as component A), a
   3-heterocyclyl-substituted benzoyl derivative of the formula
   I, where R⁴ is hydrogen.
- 5. A synergistic herbicidal mixture as claimed in any of claims
  1 to 4, comprising, as component A), a 3-heterocyclylsubstituted benzoyl derivative of the formula I, where
  - $R^1$ ,  $R^3$  are halogen,  $C_1-C_6$ -alkyl,  $C_1-C_6$ -alkylthio,  $C_1-C_6$ -alkylsulfinyl or  $C_1-C_6$ -alkylsulfonyl.
  - 6. A synergistic herbicidal mixture as claimed in any of claims 1 to 5, comprising, as component A), a

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3-heterocyclyl-substituted benzoyl derivative of the formula I, where

- is a heterocyclic radical selected from the group:
  isoxazol-3-yl, isoxazol-5-yl and
  4,5-dihydroisoxazol-3-yl, it being possible for
  the three radicals mentioned to be unsubstituted
  or mono- or polysubstituted by halogen,
  C1-C4-alkyl, C1-C4-alkoxy, C1-C4-haloalkyl,
  C1-C4-haloalkoxy or C1-C4-alkylthio.
  - 7. A synergistic herbicidal mixture as claimed in any of claims 1 to 6, comprising, as component A), a 3-heterocyclyl-substituted benzoyl derivative of the formula I, where
- is isoxazol-5-yl, 3-methyl-isoxazol-5-yl,
  4,5-dihydroisoxazol-3-yl,
  5-methyl-4,5-dihydroisoxazol-3-yl,
  5-ethyl-4,5-dihydroisoxazol-3-yl or
  4,5-dimethyl-4,5-dihydroisoxazol-3-yl.
  - 8. A synergistic herbicidal mixture as claimed in any of claims 1 to 7, comprising, as component A), 4-[2-chloro-3-(4,5-dihydroisoxazol-3-yl)-4-methylsulfonyl-benzoyl]-1-methyl-5-hydroxy-1H-pyrazole.
- A synergistic herbicidal mixture as claimed in any of claims 1 to 7, comprising, as component A),
   4-[2-methyl-3-(4,5-dihydroisoxazol-3-yl)-4-methylsulfonyl-benzoyl]-1-methyl-5-hydroxy-1H-pyrazole.
- 10. A synergistic herbicidal mixture as claimed in any of claims 1 to 5, comprising, as component A), a
  35. 3-heterocyclyl-substituted benzoyl derivative of the formula I, where
- is a heterocyclic radical selected from the group:
  thiazol-2-yl, thiazol-4-yl, thiazol-5-yl,
  isoxazol-4-yl, 4,5-dihydroisoxazol-4-yl and
  4,5-dihydroisoxazol-5-yl, it being possible for the
  six radicals mentioned to be unsubstituted or monoor polysubstituted by halogen, C₁-C₄-alkyl,
  C₁-C₄-alkoxy, C₁-C₄-haloalkyl, C₁-C₄-haloalkoxy or
  C₁-C₄-alkylthio.

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- 11. A synergistic herbicidal mixture as claimed in any of claims 1 to 10, comprising, as component B), at least one herbicidal compound from the groups B1, B2, B4 to B12 or B14 as defined in claim 2 or 3.
- 12. A synergistic herbicidal mixture as claimed in any of claims 1 to 11, comprising, as component B), at least one herbicidal compound from the following groups:
- 10 B1 acetyl-CoA carboxylase inhibitors (ACC):
  cyclohexenone oxime ethers or phenoxypropionic esters;
  - B2 acetolactate synthase inhibitors (ALS): imidazolinones, pyrimidyl ethers, sulfonamides or sulfonylureas;
  - B4 auxin herbicides: pyridinecarboxylic acids or 2,4-D;
- 20 B5 auxin transport inhibitors;
  - B6 carotenoid biosynthesis inhibitors;
  - B7 enolpyruvylshikimate 3-phosphate synthase inhibitors;

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- B8 glutamine synthetase inhibitors;
- B9 lipid biosynthesis inhibitors:
   chloroacetanilides or thioureas;

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- B10 mitosis inhibitors: dinitroanilines;
- Bll protoporphyrinogen IX oxidase inhibitors:
  diphenyl ethers, oxadiazoles, cyclic imides or pyrazoles;
  - B12 photosynthesis inhibitors: pyridate, pyridafol, benzothiadiazinone, dipyridylenes, ureas, phenols, chloridazon, triazines or triazinones;
  - B14 growth substances:

    aryloxyalkanoic acids, benzoic acids or
    quinolinecarboxylic acids.

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- 13. A synergistic herbicidal mixture as claimed in any of claims 1 to 12, comprising, as component B), at least one herbicidal compound from the following groups:
- cycloxydim, sethoxydim, clodinafop (and, if appropriate, cloquintocet), fenoxaprop-ethyl, fenoxaprop-P-ethyl, imazapyr, imazaquin, imazamethabenz, imazethapyr, pyrithiobac-sodium, metosulam, halosulfuron-methyl, nicosulfuron, N-[[[4-methoxy-6-(trifluoromethyl)-
- 1,3,5-triazin-2-yl]amino]carbonyl]-2-(trifluoromethyl)benzenesulfonamide, flufenacet, 2,4-D, diflufenzopyr,
  isoxaflutole, sulcotrione, glyphosate, glufosinate-ammonium,
  dimethenamid, S-metolachlor, benthiocarb, pendimethalin,
  acifluorfen, carfentrazone-ethyl, cinidon-ethyl, pyridate,
- bentazone, paraquat-dichloride, diuron, isoproturon, bromoxynil, chloridazon, atrazin, metribuzin, MCPA, dicamba and quinclorac.
- 14. A synergistic herbicidal mixture as claimed in any of claims

  1 to 12, comprising, as component B), at least one herbicidal compound from the group:

  codinafop (and, if appropriate, cloquintocet), diflufenzopyr, imazethapyr, flumetsulam, pyrithiobac-sodium, nicosulfuron,

  N-[[[4-methoxy-6-(trifluoromethyl)-1,3,5-triazin-2-yl]amino]carbonyl]-2-(trifluoromethyl)-benzenesulfonamide, clopyralid,
- 2,4-D, isoxaflutole, glyphosate, glufosinate-ammonium, dimethenamid, S-dimethenamid, acetochlor, metolachlor, S-metolachlor, pendimethalin, carfentrazone-ethyl, pyridate, bentazone, diuron, bromoxynil, atrazine, terbuthylazine, metribuzin and dicamba.
  - 15. A synergistic herbicidal mixture as claimed in any of claims 1 to 5, 11 to 14, comprising, as component A), a 3-heterocyclyl-substituted benzoyl derivative of the formula I where
- is a heterocyclic radical from the group:
  4,5-dihydroisoxazol-3-yl, 4,5-dihydroisoxazol-4-yl and
  4,5-dihydroisoxazol-5-yl, it being possible for the three
  radicals mentioned to be unsubstituted or mono- or
  polysubstituted by halogen, C₁-C₄-alkyl, C₁-C₄-alkoxy,
  C₁-C₄-haloalkyl, C₁-C₄-haloalkoxy or C₁-C₄-alkylthio.

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- 16. A synergistic herbicidal mixture as claimed in any of claims 1 to 5, 11 to 14, comprising, as component A), a 3-heterocyclyl-substituted benzoyl derivative of the formula I, where
- is a heterocyclic radical from the group: thiazol-2-yl, thiazol-4-yl and thiazol-5-yl, it being possible for the three radicals mentioned to be unsubstituted or mono- or polysubstituted by halogen, C₁-C₄-alkyl, C₁-C₄-alkoxy, C₁-C₄-haloalkyl, C₁-C₄-haloalkyl or C₁-C₄-alkylthio.
- 17. A synergistic herbicidal mixture as claimed in any of claims 1 to 5, 11 to 14, comprising, as component A), a 3-heterocyclyl-substituted benzoyl derivative of the formula I, where
- is a heterocyclic radical from the group: isoxazol-3-yl, isoxazol-4-yl or isoxazol-5-yl, it being possible for the three radicals mentioned to be unsubstituted or mono- or polysubstituted by halogen, C₁-C₄-alkyl, C₁-C₄-alkoxy, C₁-C₄-haloalkyl, C₁-C₄-haloalkoxy or C₁-C₄-alkylthio.
- 18. A synergistic herbicidal mixture as claimed in any of claims 15 to 17, comprising, as component B), at least one herbicidal compound from the groups B1, B2, B4 to B11 or B14 as defined in claim 2.
- 19. A synergistic herbicidal mixture as claimed in claim 15 or 16, comprising, as component B), at least one herbicidal
  30 compound from the groups B12 as defined in claim 2.
  - 20. A synergistic herbicidal mixture as claimed in any of claims 15 to 17, comprising, as component B), at least one herbicidal compound from the following group:
    - propanil, pyridate, pyridafol, dinitrophenols, dipyridylenes, triazinones, uracils or biscarbamates.
- 21. A synergistic herbicidal mixture as claimed in any of claims 1 to 20, comprising, as component A), a 3-heterocyclyl-substituted benzoyl derivative of the formula I and, as component B), a herbicidal compound as defined in any of claims 1 to 20.

- 22. A synergistic herbicidal mixture as claimed in any of claims 1 to 20, comprising, as component A), a 3-heterocyclyl-
- substituted benzoyl derivative of the formula I and, as component B), two herbicidal compounds as defined in any of claims 1 to 20.
- 23. A synergistic herbicidal mixture as claimed in any of claims 1 to 12, comprising a 3-heterocyclyl-substituted
  10 benzoyl derivative of the formula I and, as component B), a herbicidal compound as defined in any of claims 1 to 12 and a herbicidal compound from the groups B12 and B14.
- 24. Synergistic herbicidal mixture as claimed in any of claims 1 to 23, wherein component A) and B) are present in a weight ratio of 1:0.002 to 1:800.
- 25. Synergistic herbicidal mixture as claimed in claim 24, wherein component A) and component B) are present in a weight ratio of 1:0.003 to 1:160.
- 26. A herbicidal composition comprising a herbicidally active amount of a synergistic herbicidal mixture as claimed in any of claims 1 to 23, at least one inert liquid and/or solid carrier and, if desired, at least one surfactant.
  - 27. A herbicidal composition as claimed in claim 26, wherein component A) and component B) are present in a weight ratio of 1:0.002 to 1:800.
  - 28. A herbicidal composition as claimed in claim 27, wherein component A) and component B) are present in a weight ratio of 1:0.003 to 1:160.
- 35 29. A process for the preparation of herbicidal compositions as claimed in claim 25, wherein component A, component B, at least one inert liquid and/or solid carrier and, if appropriate, a surfactant are mixed.
- 40 30. A method of controlling undesired vegetation, which comprises applying a synergistic herbicidal mixture as claimed in any of claims 1 to 23 before, during and/or after the emergence of undesired plants, it being possible for the herbicidally active compounds of components A) and B) to be applied simultaneously or in succession.

31. A method of controlling undesired vegetation as claimed in claim 30, wherein the leaves of the crop plants and of the undesired plants are treated.

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